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INTERN'ATIONAL ADDITICATION DIDLICUED HINDED THE DATENT COODED ATION T

51) International Patent Classification ⁶ :			(11) International Publication Number: WO 99/5713
C07K 14/00		A2	(43) International Publication Date: 11 November 1999 (11.11.99
21) International Application Number:	PCT/US	99/0952	(74) Agents: TSAO, Y., Rocky et al., Fish & Richardson P.C., 22 Franklin Street, Boston, MA 02110-2804 (US).
22) International Filing Date: 3 M	May 1999 (03.05.9	
30) Priority Data: 09/072,956 5 May 1998 (05.0) 63) Related by Continuation (CON) or Conti (CIP) to Earlier Application US Filed on 5 M 71) Applicant (for all designated States except CONSEILS DE RECHERCHES ET SCIENTIFIQUES S.A. [FR/FR]; 51-5 Blanche, F-75016 Paris (FR). 72) Inventors; and 75) Inventors/Applicants (for US only): [IL/US]; 205 South Street, Chestnut (US). DONG, Zheng, Xin [CN/US]; Framingham, MA 01701 (US). ROSE [US/US]; 130 Lake Avenue, Newton (US).	inuation-ir 09/072,95 May 1998 (i US): SOC D'APPLIC 53, rue du CHOREV, t Hill, M. 40 Angelic NBLATT,	Micha A 0216 Driv Micha Micha Micha Micha Micha Micha	KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, M MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, S SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, Z ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SL, S UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, M RU, TJ, TM), European patent (AT, BE, CH, CY, DE, D ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OA patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, M NE, SN, TD, TG). Published Without international search report and to be republish upon receipt of that report.
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WO 99/57139 PCT/US99/09521

- 1 -

PTH2 RECEPTOR SELECTIVE COMPOUNDS Statement as to Government Funding

This invention was supported in part by Government funding, NIDDK Research Grant DK-4790, and the Government, therefore, may have certain rights in the invention.

Background of the Art

This invention relates to a series of PTH and PTHrP analogues that selectively bind to PTH2 receptors and as such may be useful in treating abnormal CNS functions; abnormal pancreatic functions; divergence from normal mineral metabolism and homeostasis; male infertility; regulation of abnormal blood pressure; and hypothalmic disease, to name a few potential uses.

An alternate human parathyroid hormone receptor, designated as PTH2 receptor, has been identified in rat and human brain. This receptor is selectively activated by PTH-(1-34), but not PTH-related protein PTHrP-20 (1-34), which has the same calcium-mobilizing activities as PTH-(1-34). Both PTH and PTHrP share a common G proteincoupled receptor, termed the PTH/PTHrP receptor. The PTH2 receptor is localized predominantly in the brain and pancreas, in contrast to PTH/PTHrP receptor, which is 25 primarily localized in bone and the kidney, the principal target tissue for PTH action. Parathyroid hormone (PTH) is the principal physiological regulator of calcium levels in the blood (Chorev, M., Rosenblatt, M., 1994, Structure function analysis of parathyroid hormone and parathyroid 30 hormone-related protein, Bilezikian, J.P., Marcus, R., Levine, M., (eds) The Parathyroids: Basic and Clinical Concepts. Raven Press, New York, pp 139-156; Juppner, H., et al., 1991, Science, 254:1024-1026; and Martin, T.J., et al., 1991, Crit. Rev. Biochem. Mol. Biol. 26:377-395).

PTH-related protein (PTHrP) was originally identified as the agent responsible for the paraneoplastic syndrome of humoral hypercalcemia of malignancy (Suva, L.J., et al., 1987, Science, 237:893-896 and Orloff, J.J., et al., 1994, 5 Endocrinol. Rev. 15:40-60). PTH and PTHrP are products of distinct, yet evolutionary-related genes. PTH and PTHrP show sequence similarities only in the N-terminal 13 amino acids, 8 of which are identical (Abou-Samra AB, et al., 1992, Proc. Natl. Sci. Acad. USA, 89:2732-2736). However, 10 the expression pattern and physiological role of these two molecules are remarkably different. PTH has a highly restricted pattern of expression and acts as a classical endocrine hormone, whereas PTHrP is expressed in a wide variety of normal tissues and functions in a predominantly 15 autocrine/paracrine fashion (Urena, P., et al., Endocrinology, 133:617-623; Lee, K., et al., Endocrinology, 136:453-463; and Martin, T.J., et al., 1995, Miner. Electrolyte Metab., 21:123-128). recently, PTHrP has been shown to play a fundamental role 20 in embryonic differentiation of bone and cartilage development.

PTH and PTHrP exert their wide-ranging effects via a common receptor located on the surface of target cells (Juppner, H., et al., 1988, J. Biol. Chem., 263:1071-1078; Shigeno, C., et al., 1988, J. Biol. Chem., 263:18369-25 18377). The PTH/PTHrP receptor is a member of a subfamily of G protein-coupled receptor superfamily, which includes the receptors for glucagon, growth hormone-releasing hormone (GHRH), vasoactive intestinal peptide (VIP), 30 glucagon-like peptide 1 (GLP-1), gastric inhibitory polypeptide (GIP), secretin, pituitary adenylate cyclaseactivating polypeptide (PACAP), calcitonin, corticotropin-releasing factor (CRF) (Segre, G., et al., 1993, Trends Endocrinol. Metab. 4:309-314). The PTH/PTHrP receptor recognizes the N-terminal 1-34 regions of both 35 ligands (Schipani, E., et al., 1993, Endocrinology, 132:2157-2165) and is particularly abundant in classical PTH target tissues such as bone and kidney (Urena, P., et

al., 1993 Endocrinology, 133:35-38). Ligand binding to the PTH/PTHrP receptor can activate at least two signaling pathways; the adenylyl cyclase-cAMP-protein kinase A pathway (Partridge, NC, et al., 1981, Endocrinology 108:220-225), and the inositol trisphosphate-cytosolic calcium-protein kinase C pathway (Abou-Samra, A-B., et al., 1989, Endocrinology 124:1107-1113).

An homologous receptor for PTH, designated the PTH2 receptor, has been identified and partially characterized 10 (Behar, V., et al., 1996, Endocrinology, 137:2748-2757; Gardella, T.J., et al., 1996, The J. Biol. 271:19888-19893; Behar, V., et al., 1996, Endocrinology, 137:4217-4224; and Usdin, T.B., et al., 138:831-834). Endocrinology, Amongst the seven 15 transmembrane G protein-coupled receptors, the PTH2 receptor is most similar in sequence to the PTH/PTHrP receptor (51% of the amino acid sequence identify). Interestingly, PTH2 receptor mRNA is not detected in bone or osteosarcoma cell lines, but is expressed in a number of 20 tissues including the exocrine pancreas, lung, heart, vasculature, and epididymis, and is most abundant in the brain (Usdin, T.B., et al., 1996, Endocrinology, 137:4285-4297). Unlike the PTH/PTHrP receptor, which binds and is activated by both PTH-(1-34) and PTHrP-(1-34), the PTH2 receptor binds and is activated only by PTH-(1-34). 25 PTHrP(7-34) was found to recognize PTH2 receptor and weakly activate it. Moreover, His⁵ in PTHrP was identified as the "specificity switch" for the PTH2 receptor. Swapping a single amino acid, His5 from PTHrP, with Ile5 from PTH, 30 resulted in a PTHrP analogue, Ile⁵-PTHrP-(1-34)NH₂, which acts as a PTH-2 receptor agonist. Hence, the single amino acid switch converts inactive PTHrP into a potent PTH2 receptor agonist. But while [Ile⁵] PTHrP binds and activates both receptors, PTH/PTHrP and PTH2, it is not a selective 35 PTH2 agonist. In transient heterologous (with respect to species) expression systems, others have found additional contribution to hPTH2 receptor selectivity by Trp²³ (Gardella et al., JBC 1996, 271:19888-19893). Like the

PTH/PTHrP receptor, PTH binding leads to PTH2 receptormediated activation of both cAMP and [Ca2+] intracellular signaling pathways.

The physiological function of the PTH2 receptor 5 because of its high abundance and distribution in the brain suggests that it may act as a neurotransmitter receptor. PTH has been found in the central nervous system (CNS) (Harvey, S., et al., 1993, J. Endocrinol. 139:353-361), therefore, it is possible that endogenous PTH2 receptor 10 specific ligands, which are distinct from PTH, do exist in Recently, Usdin reported the isolation of "PTH2 receptor binding activity" from the hypothalamus which was immunologically distinct from PTH.

PCT Application Number PCT/US97/13360, published as PCT Publication Number WO 98/04591, discloses the use of certain PTHrP analogs which are PTH2 receptor agonists or antagonists.

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U.S. Patent No. 5,723,577, issued March 3, 1998, discloses certain PTH and PTHrP analogues. U.S. Application 20 Nos. 08/779,768 and 08/813,534, filed January 7, 1997 and March 7, 1997, respectively, disclose further PTH and PTHrP analogs.

The development of specific ligands which activate the PTH2 receptor but not the PTH/PTHrP receptor, would be 25 highly useful in defining the physiological roles of the PTH2 receptor and its potential involvement in certain pathological states. We have discovered a series of PTH2 receptor-selective PTH analogues which interact selectively with the human PTH2 receptor and are practically devoid of 30 PTH/PTHrP receptor interaction. The compounds of the present invention are not only selective toward a receptor subtype but also signal specifically through stimulation of [Ca⁺²], transients. Therefore, the compounds of the present invention are receptor subtype and signaling pathway selective.

Summary of the Invention

In one aspect, this invention provides a analogue or a truncated PTH analogue or a pharmaceutically acceptable salt thereof that selectively binds to the PTH2 receptor. A preferred PTH analogue or a truncated PTH analogue or a pharmaceutically acceptable salt thereof is where the analogue is a selective PTH2 receptor agonist.

5 Another preferred PTH analogue or a truncated PTH analogue or a pharmaceutically acceptable salt thereof is where the analogue is a selective PTH2 receptor antagonist.

A more preferred PTH analogue that selectively binds to the PTH2 receptor is an analogue of formula (I),

10 $(R^1R^2) - A^1 - A^2 - A^3 - A^4 - A^5 - A^6 - A^7 - A^8 - A^9 - A^{10} - A^{11} - A^{12} - A^{13} - A^{14} - A^{15} - A^{16} - A^{17} - A^{18} - A^{19} - A^{20} - A^{21} - A^{22} - A^{23} - A^{24} - A^{25} - A^{26} - A^{27} - A^{28} - A^{29} - A^{30} - A^{31} - A^{32} - A^{33} - A^{34} - A^{35} - A^{36} - A^{37} - A^{38} - R^3$

(I)

or a pharmaceutically-acceptable salt thereof wherein

15 A' is a hydrophilic or a lipophilic amino acid;

A² is a lipophilic amino acid;

A³ is a hydrophilic or a lipophilic amino acid;

A4 is a hydrophilic amino acid;

A⁵ is a hydrophilic or a lipophilic amino acid;

20 A^6 is a hydrophilic amino acid or is deleted; A^7 is a hydrophilic or a lipophilic amino acid or is deleted;

A⁸ is a lipophilic amino acid or is deleted;

A9 is a hydrophilic amino acid or is deleted;

25 A¹⁰ is a hydrophilic amino acid or is deleted;

A¹¹ is a hydrophilic or a lipophilic amino acid or is deleted;

A¹² is a hydrophilic or a lipophilic amino acid or is deleted;

30 A¹³ is a hydrophilic amino acid;

A14 is a hydrophilic amino acid or is deleted;

A¹⁵ is a lipophilic amino acid or is deleted;

A¹⁶ is a hydrophilic or a lipophilic amino acid or is deleted;

35 A¹⁷ is a hydrophilic or a lipophilic amino acid or is deleted;

A¹⁸ is a lipophilic amino acid or is deleted;

A¹⁹ is a hydrophilic or a lipophilic amino acid or is deleted;

A²⁰ is a hydrophilic amino acid or is deleted;

 A^{21} is a hydrophilic or a lipophilic amino acid or is deleted;

A²² is a lipophilic or a hydrophilic amino acid or is deleted;

A²³ is a hydrophilic or a lipophilic amino acid;

A²⁴ is a hydrophilic or a lipophilic amino acid;

10 A²⁵ is a hydrophilic amino acid;

A²⁶ is a hydrophilic amino acid;

A²⁷ is a lipophilic or a hydrophilic amino acid;

A²⁸ is a lipophilic amino acid;

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A²⁹ is a lipophilic or a hydrophilic amino acid;

15 A^{30} is a hydrophilic or a lipophilic amino acid;

A³¹ is a lipophilic or a hydrophilic amino acid or is deleted;

A³² is a hydrophilic amino acid or is deleted;

A³³ is a hydrophilic amino acid or is deleted;

20 A³⁴ is a lipophilic amino acid or is deleted;

A³⁵ is a lipophilic amino acid or is deleted;

 A^{36} is a lipophilic or a hydrophilic amino acid or is deleted;

A³⁷ is a lipophilic amino acid or is deleted;

25 A³⁸ is a lipophilic or a hydrophilic amino acid or is deleted;

 R^1 and R^2 are each independently selected from the group consisting of H, (C_{1-30}) alkyl, (C_{2-30}) alkenyl, phenyl- (C_{1-30}) alkyl, naphthyl (C_{1-30}) alkyl, hydroxy (C_{1-30}) alkyl, hydroxy (C_{2-30}) alkenyl, hydroxy-phenyl (C_{1-30}) alkyl or hydroxy-naphthyl (C_{1-30}) alkyl; or one of R^1 or R^2 is COE^1 where E^1 is (C_{1-30}) alkyl, (C_{2-30}) alkenyl, phenyl (C_{1-30}) alkyl, naphthyl (C_{1-30}) alkyl, hydroxy (C_{1-30}) alkyl, hydroxy (C_{2-30}) alkenyl, hydroxy-phenyl (C_{1-30}) alkyl or hydroxy-naphthyl (C_{1-30}) alkyl; and R^3 is OH, NH_2 , (C_{1-30}) alkoxy or $NH-Y-CH_2-Z$, where Y is

R³ is OH, NH₂, (C_{1-30}) alkoxy or NH-Y-CH₂-Z, where Y is a (C_{1-30}) hydrocarbon moiety and Z is CO_2H or $CONH_2$;

provided that the compound is not $PTH(1-34)R^3$, $PTH(1-35)R^3$, $PTH(1-36)R^3$, $PTH(1-37)R^3$, or $PTH(1-38)R^3$.

Another preferred group of PTH analogues that selectively binds to the PTH2 receptor is an analogue of formula (II),

 $(R^1R^2) - A^1 - A^2 - A^3 - A^4 - A^5 - A^6 - A^7 - A^8 - A^9 - A^{10} - A^{11} - A^{12} - A^{13} - A^{14} - A^{15} - A^{16} - A^{17} - A^{18} - A^{19} - A^{20} - A^{21} - A^{22} - A^{23} - A^{24} - A^{25} - A^{26} - A^{27} - A^{28} - A^{29} - A^{30} - A^{31} - A^{32} - A^{33} - A^{34} - A^{35} - A^{36} - A^{37} - A^{38} - R^3 ,$

(II)

or a pharmaceutically-acceptable salt thereof wherein A^1 is Ser, Ala, Dap, Thr, Aib or is deleted; A^2 is Val, Leu, Ile, Phe, Nle, β -Nal, Aib, p-X-Phe, Acc, Cha, Met or is deleted; A^3 is Ser, Thr, Aib or is deleted;

15 A^4 is Glu, Asp or is deleted; A^5 is Leu, Val, Nle, Ile, Cha, β -Nal, Trp, Pal, Acc, Phe, p-X-Phe or is deleted; A^6 is Gln, a hydrophilic amino acid or is deleted;

 A^7 is Leu, Val, Nle, Ile, Cha, β -Nal, Trp, Pal, Acc, Phe, p-20 X-Phe, a lipophilic amino acid, or is deleted; A^8 is Met, Nva, Leu, Val, Ile, Cha, Acc, Nle, p-X-Phe, Phe, β -Nal, Bpa, a lipophilic amino acid or is deleted; A^9 is His, a hydrophilic amino acid or is deleted;

A¹⁰ is Asn, a hydrophilic amino acid or is deleted;

25 A¹¹ is Leu, Val, Nle, Ile, Cha, β -Nal, Trp, Pal, Acc, Phe, p-X-Phe, a hydrophilic amino acid or is deleted; A¹² is Gly, Acc, Aib, or is deleted; A¹³ is Lys, Arg or HN-CH((CH₂)_nNH-R⁴)-C(O);

A¹⁴ is His or is deleted;

30 A^{15} is Leu, Val, Nle, Ile, Cha, β -Nal, Trp, Pal, Acc, Phe, p-X-Phe or is deleted;

A¹⁶ is Ser, Asn, Ala, Aib or is deleted;

A¹⁷ is Ser, Thr, Aib or is deleted;

 A^{18} is Met, Nva, Leu, Val, Ile, Nle, p-X-Phe, Phe, β -Nal,

35 Acc, Cha, Aib or is deleted;

A19 is Glu, Aib or is deleted;

 A^{20} is Arq, Lys, HN-CH((CH₂),NH-R⁴)-C(0) or is deleted;

 A^{21} is Val, Leu, Ile, Phe, Nle, β -Nal, Aib, p-X-Phe, Acc, Cha, Met or is deleted;

A²² is Acc, Aib, Glu or is deleted;

 A^{23} is Trp, Acc, Phe, p-X-Phe, Aib, β -Nal or Cha;

5 A²⁴ is Leu, Acc, Ile, Val, Phe, β -Nal, Nle, Aib, p-X-Phe or Cha;

 A^{25} is Arg, Lys or $HN-CH((CH_2)_nNH-R^4)-C(0)$;

 A^{26} is Arg, Lys or $HN-CH((CH_2)_nNH-R^4)-C(O)$;

A²⁷ is Lys, Aib, Leu, hArg, Gln, Acc, Arg, Cha, Nle, Ile,

10 Val, Phe, β -Nal, or p-X-Phe, where the Lys is optionally substituted on the ϵ -amino group by an acyl group; A²⁸ is Leu, Acc, Cha, Ile, Val, Phe, Nle, β -Nal, Aib or p-X-Phe;

A²⁹ is Gln, Acc or Aib;

15 A³⁰ is Asp, Lys, Arg or is deleted;

 A^{31} is Val, Leu, Nle, Acc, Cha, Phe, Ile, β -Nal Aib, p-X-Phe or is deleted;

A³² is His or is deleted;

A33 is Asn or is deleted;

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20 A^{34} is Phe, Tyr, Amp, Aib, β -Nal, Cha, Nle, Leu, Ile, Acc, p-X-Phe or is deleted;

 A^{35} is Val, Leu, Nle, Acc, Cha, Phe, Ile, β -Nal Aib, p-X-Phe or is deleted;

A³⁶ is Ala, Val, Aib, Acc, Nva, Abu or is deleted;

25 A^{37} is Leu, Val, Nle, Ile, Cha, β -Nal, Trp, Pal, Acc, Phe, p-X-Phe, a lipophilic amino acid, or is deleted; A^{38} is Gly, Acc, Aib, or is deleted;

where X for each occurrence is independently selected from the group consisting of OH, a halo and CH₃;

 R^1 and R^2 are each independently selected from the group consisting of H, (C_{1-30}) alkyl, (C_{2-30}) alkenyl, phenyl- (C_{1-30}) alkyl, naphthyl (C_{1-30}) alkyl, hydroxy (C_{1-30}) alkyl, hydroxy (C_{2-30}) alkenyl, hydroxy-phenyl (C_{1-30}) alkyl or hydroxy-naphthyl (C_{1-30}) alkyl;

30) alkyl or hydroxy-naphthyl (C_{1-30}) alkyl; or one of R^1 or R^2 is COE^1 where E^1 is (C_{1-30}) alkyl, (C_{2-30}) alkenyl, phenyl (C_{1-30}) alkyl, naphthyl (C_{1-30}) alkyl, hydroxy (C_{1-30}) alkyl, hydroxy (C_{2-30}) alkenyl,

hydroxy-phenyl (C_{1-30}) alkyl or hydroxy-naphthyl (C_{1-30}) alkyl;

 R^3 is OH, NH_2 , (C_{1-30}) alkoxy or $NH-Y-CH_2-Z$, where Y is a (C_{1-30}) hydrocarbon moiety and Z is CO_2H or $CONH_2$; n for each occurrence is independently an integer from 1 to 5; and

 R^4 for each occurrence is independently (C_1-C_{30}) alkyl, (C_1-C_{30}) acyl or $-C((NH)(NH_2))$;

provided that the compound is not $PTH(1-34)R^3$, $PTH(1-35)R^3$, 10 $PTH(1-36)R^3$, $PTH(1-37)R^3$, or $PTH(1-38)R^3$.

5

In another aspect, this invention provides a PTHrP analogue that selectively binds to the PTH2 receptor of the formula (IV),

 $(R^1R^2) - A^1 - A^2 - A^3 - A^4 - A^5 - A^6 - A^7 - A^8 - A^9 - A^{10} - A^{11} - A^{12} - A^{13} - A^{14} - A^{15} - A^{16} - A^{17} - A^{18} -$

15 $A^{19} - A^{20} - A^{21} - A^{22} - A^{23} - A^{24} - A^{25} - A^{26} - A^{27} - A^{28} - A^{29} - A^{30} - A^{31} - A^{32} - A^{33} - A^{34} - A^{35} - A^{36} - A^{37} - A^{38} - R^{3}$,

(IV)

or a pharmaceutically acceptable salt thereof, wherein ${\tt A^1}$ is Ala, Ser, Dap, Thr, Aib or is deleted;

- 20 A² is Val or is deleted; A³ is Ser, Aib, Thr or is deleted; A⁴ is Glu, Asp or is deleted; A⁵ is His, Ile, Acc, Val, Nle, Phe, Leu, p-X-Phe, β -Nal, Aib, Cha or is deleted;
- 25 A⁶ is Gln, a hydrophilic amino acid or is deleted; A⁷ is Leu, Val, Cha, Nle, β -Nal, Trp, Pal, Acc, Phe, p-X-Phe, Aib, a lipophilic amino acid or is deleted; A⁸ is Leu, Met, Acc, Cha, Aib, Nle, Phe, Ile, Val, β -Nal, p-X-Phe, a lipophilic amino acid or is deleted;
- 30 A° is His, a hydrophilic amino acid or is deleted; A¹⁰ is Asp, Asn, a hydrophilic amino acid or is deleted; A¹¹ is Lys, Arg, Leu, Cha, Aib, p-X-Phe, Ile, Val, Nle, Acc, Phe, β -Nal, HN-CH((CH₂) $_n$ NH-R⁴)-C(O), a lipophilic D-amino acid, a hydrophilic amino acid or is deleted;
- 35 A^{12} is Gly, Acc, Aib or is deleted; A^{13} is Lys, Arg, HN-CH((CH₂)_nNH-R⁴)-C(0) or is deleted; A^{14} is Ser, His or is deleted;

 A^{15} is Ile, Acc, Cha, Leu, Phe, Nle, β -Nal, Trp, p-X-Phe, Val, Aib or is deleted;

A¹⁶ is Gln, Aib or is deleted;

A¹⁷ is Asp, Aib or is deleted;

5 A^{18} is Leu, Aib, Acc, Cha, Phe, Ile, Nle, β -Nal, Val, p-X-Phe or is deleted;

 A^{19} is Arg, Lys, Aib, $HN-CH((CH_2)_nNH-R^4)-C(0)$ or is deleted; A^{20} is Arg, Lys, $HN-CH((CH_2)_nNH-R^4)-C(0)$ or is deleted;

 A^{21} is Arg, Lys, $HN-CH((CH_2)_nNH-R^4)-C(0)$ or is deleted;

10 A^{22} is Phe, Glu, Aib, Acc, p-X-Phe, β -Nal, Val, Leu, Ile, Nle or Cha;

 A^{23} is Phe, Leu, Lys, Acc, Cha, β -Nal, Aib, Nle, Ile, p-X-Phe, Val or Trp;

 A^{24} is Leu, Lys, Acc, Nle, Ile, Val, Phe, β -Nal, Aib, p-X-

15 Phe, Arg or Cha;

A²⁵ is His, Lys, Aib, Acc, Arg or Glu;

A²⁶ is His, Aib, Acc, Arg or Lys;

 ${\rm A^{27}}$ is Leu, Lys, Acc, Arg, Ile, Val, Phe, Aib, Nle, ${\it \beta}$ -Nal, p-X-Phe or Cha;

20 A^{28} is Ile, Leu, Lys, Acc, Cha, Val, Phe, p-X-Phe, Nle, β -Nal, Aib or is deleted;

A²⁹ is Ala, Glu, Acc, Aib or is deleted;

 A^{30} is Glu, Leu, Nle, Cha, Aib, Acc, Lys, Arg or is deleted; A^{31} is Ile, Leu, Cha, Lys, Acc, Phe, Val, Nle, β -Nal, Arg or

25 is deleted;

A³² is His or is deleted;

A³³ is Thr, Ser or is deleted;

 A^{34} is Ala, Phe, Tyr, Cha, Val, Ile, Leu, Nle, β -Nal, Aib, Acc or is deleted;

30 A³⁵ is Glu, Asp or is deleted;

 A^{36} is Ile, Acc, Cha, Leu, Phe, Nle, β -Nal, Trp, p-X-Phe, Val, Aib or is deleted;

 A^{37} is Arg, Lys, $HN-CH((CH_2)_nNH-R^4)-C(0)$ or is deleted;

 A^{38} is Ala, Phe, Tyr, Cha, Val, Ile, Leu, Nle, β -Nal, Aib,

35 Acc or is deleted;

 R^1 and R^2 are each independently selected from the group consisting of H, (C_{1-30}) alkyl, (C_{2-30}) alkenyl, phenyl- (C_{1-30}) alkyl, naphthyl (C_{1-30}) alkyl, hydroxy (C_{1-30})

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alkyl, hydroxy(C_{2-30}) alkenyl, hydroxy-phenyl(C_{1-30}) alkyl or hydroxy-naphthyl(C_{1-30}) alkyl; or one of R^1 or R^2 is COE^1 where E^1 is (C_{1-30}) alkyl, (C_{2-30}) alkenyl, phenyl(C_{1-30}) alkyl, naphthyl(C_{1-30}) alkyl, hydroxy(C_{2-30}) alkenyl, hydroxy(C_{2-30}) alkyl, hydroxy-phenyl(C_{1-30}) alkyl or hydroxy-naphthyl(C_{1-30}) alkyl;

 R^3 is OH, NH_2 , (C_{1-30}) alkoxy or $NH-Y-CH_2-Z$, where Y is a (C_{1-30}) hydrocarbon moiety and Z is CO_2H or $CONH_2$; n for each occurrence is independently an integer from 1 to 5; and

 R^4 for each occurrence is independently (C_1 - C_{30}) alkyl, (C_1 - C_{30}) acyl or -C((NH)(NH₂));

provided that the compound is not PTHrP(1-34)R³, PTHrP(1-35)R³, PTHrP(1-36)R³, PTHrP(1-37)R³ or PTHrP(1-38)R³, and further provided that the compound is not [Ile⁵, Trp²³] PTHrP(1-36) or [Trp²³] PTHrP(1-36).

In another aspect, this invention provides a method of selectively binding the PTH2 receptor which comprises administering to a patient in need thereof an effective amount of a PTH analogue or a truncated PTH analogue or a pharmaceutically acceptable salt thereof that selectively binds to a PTH2 receptor.

In another aspect, this invention provides a method of selectively eliciting an agonist response from the PTH2 receptor which comprises administering to a patient in need thereof an effective amount of a PTH analogue or a truncated PTH analogue or a pharmaceutically acceptable salt thereof which is a selective PTH2 receptor agonist.

In another aspect, this invention provides a method of selectively eliciting an antagonist response from the PTH2 receptor which comprises administering to a patient in need thereof an effective amount of a PTH analogue or a truncated PTH analogue or a pharmaceutically acceptable salt thereof which is a selective PTH2 receptor antagonist.

In yet another aspect, this invention provides a compound of the formula (III),

 $\begin{array}{l} (R^1R^2) - A^1 - A^2 - A^3 - A^4 - A^5 - A^6 - A^7 - A^8 - A^9 - A^{10} - A^{11} - A^{12} - A^{13} - A^{14} - A^{15} - A^{16} - A^{17} - A^{18} - A^{19} - A^{20} - A^{21} - A^{22} - A^{23} - A^{24} - A^{25} - A^{26} - A^{27} - A^{26} - A^{29} - A^{30} - A^{31} - A^{32} - A^{33} - A^{34} - A^{35} - A^{36} - A^{37} - A^{38} - R^3 \,, \end{array}$

(III)

or a pharmaceutically-acceptable salt thereof wherein A^1 is Ser, Ala, Dap, Thr, Aib or is deleted; A^2 is Val, Leu, Ile, Phe, Nle, β -Nal, Aib, p-X-Phe, Acc, Cha, Met or is deleted;

A3 is Ser, Thr, Aib or is deleted;

10 A 4 is Glu, Asp or is deleted; A 5 is Leu, Val, Nle, Ile, Cha, β -Nal, Trp, Pal, Acc, Phe, p-X-Phe or is deleted;

 A^6 is Gln, a hydrophilic amino acid or is deleted; A^7 is Leu, Val, Nle, Ile, Cha, β -Nal, Trp, Pal, Acc, Phe, p-

- 15 X-Phe, a lipophilic amino acid, or is deleted;
 A⁸ is Met, Nva, Leu, Val, Ile, Cha, Acc, Nle, p-X-Phe, Phe,
 β-Nal, Bpa, a lipophilic amino acid or is deleted;
 A⁹ is His, a hydrophilic amino acid or is deleted;
 A¹⁰ is Asn, a hydrophilic amino acid or is deleted;
- 20 A¹¹ is Leu, Val, Nle, Ile, Cha, β-Nal, Trp, Pal, Acc, Phe, p-X-Phe, a hydrophilic amino acid or is deleted; A¹² is Gly, Acc, Aib, or is deleted; A¹³ is Lys, Arg or HN-CH((CH₂)_nNH-R⁴)-C(O); A¹⁴ is His or is deleted;
- 25 A^{15} is Leu, Val, Nle, Ile, Cha, β -Nal, Trp, Pal, Acc, Phe, p-X-Phe or is deleted; A^{16} is Ser, Asn, Ala, Aib or is deleted; A^{17} is Ser, Thr, Aib or is deleted; A^{18} is Met, Nva, Leu, Val, Ile, Nle, p-X-Phe, Phe, β -Nal,
- 30 Acc, Cha, Aib or is deleted; $A^{19} \text{ is Glu, Aib or is deleted;} \\ A^{20} \text{ is Arg, Lys, HN-CH((CH_2)_nNH-R^4)-C(0) or is deleted;} \\ A^{21} \text{ is Val, Leu, Ile, Phe, Nle, β-Nal, Aib, p-X-Phe, Acc, Cha, Met or is deleted;} \\$
- 35 A^{22} is Acc, Aib, Glu or is deleted; A^{23} is Trp, Acc, Phe, p-X-Phe, Aib, β -Nal or Cha; A^{24} is Leu, Acc, Ile, Val, Phe, β -Nal, Nle, Aib, p-X-Phe or Cha;

 A^{25} is Arg, Lys or $HN-CH((CH_2)_nNH-R^4)-C(O)$;

 A^{26} is Arg, Lys or HN-CH((CH₂)_NH-R⁴)-C(O);

 A^{27} is Lys, Aib, Leu, hArg, Gln, Acc, Arg, Cha, Nle, Ile, Val, Phe, β -Nal, or p-X-Phe, where the Lys is optionally

5 substituted on the ϵ -amino group by an acyl group; A^{28} is Leu, Acc, Cha, Ile, Val, Phe, Nle, β -Nal, Aib or p-X-Phe;

A²⁹ is Gln, Acc or Aib;

A³⁰ is Asp, Lys, Arg or is deleted;

10 A^{31} is Val, Leu, Nle, Acc, Cha, Phe, Ile, β -Nal Aib, p-X-Phe or is deleted;

A³² is His or is deleted;

A33 is Asn or is deleted;

 A^{34} is Phe, Tyr, Amp, Aib, β -Nal, Cha, Nle, Leu, Ile, Acc,

15 p-X-Phe or is deleted;

 A^{35} is Val, Leu, Nle, Acc, Cha, Phe, Ile, β -Nal Aib, p-X-Phe or is deleted;

A³⁶ is Ala, Val, Aib, Acc, Nva, Abu or is deleted;

 A^{37} is Leu, Val, Nle, Ile, Cha, β -Nal, Trp, Pal, Acc, Phe,

20 p-X-Phe, a lipophilic amino acid, or is deleted; A³⁸ is Gly, Acc, Aib, or is deleted;

where X for each occurrence is independently selected from the group consisting of OH, a halo and CH₃;

- 25 R¹ and R² are each independently selected from the group consisting of H, (C_{1-30}) alkyl, (C_{2-30}) alkenyl, phenyl- (C_{1-30}) alkyl, naphthyl (C_{1-30}) alkyl, hydroxy (C_{1-30}) alkyl, hydroxy (C_{2-30}) alkenyl, hydroxy-phenyl (C_{1-30}) alkyl or hydroxy-naphthyl (C_{1-30}) alkyl;
- or one of R^1 or R^2 is COE^1 where E^1 is (C_{1-30}) alkyl, (C_{2-30}) alkenyl, phenyl (C_{1-30}) alkyl, naphthyl (C_{1-30}) alkyl, hydroxy (C_{1-30}) alkyl, hydroxy (C_{2-30}) alkenyl, hydroxy-phenyl (C_{1-30}) alkyl or hydroxy-naphthyl (C_{1-30}) alkyl;
- R³ is OH, NH₂, (C_{1-30}) alkoxy or NH-Y-CH₂-Z, where Y is a (C_{1-30}) hydrocarbon moiety and Z is CO_2H or $CONH_2$; n for each occurrence is independently an integer from 1 to 5; and

 R^4 for each occurrence is independently (C₁-C₃₀)alkyl, (C₁-C₃₀)acyl or -C((NH)(NH₂));

provided that when A⁸ is not a lipophilic D-amino acid or is not deleted then at least one of A⁶, A⁷, A⁹, A¹⁰, A¹¹ and A¹² is a D-amino acid or at least one of A⁶, A⁷, A⁹, A¹⁰, A¹¹, A¹², A¹³, A¹⁴, A¹⁵, A¹⁶, A¹⁷, A¹⁸, A¹⁹, A²⁰, A²¹ and A²² is deleted; and further provided that when the compound contains a D-amino acid then A³⁶ is deleted.

A preferred group of compounds of formula (III) are the compounds listed as Examples 1-73, shown hereinbelow. Of the compounds listed as Examples 1-73, the following compounds are preferred: [Cha^{7,11}, des-Met⁸, Nle¹⁸, Tyr³⁴]hPTH-(1-34)NH₂,

[Cha^{7,11}, D-Nle⁸, des-Met¹⁸, Tyr³⁴] hPTH-(1-34)NH₂, [Cha^{7,11}, D-15 Nle⁸, Nle¹⁸, Tyr³⁴] hPTH-(1-34)NH₂, [D-Nle⁸, Nle¹⁸, Tyr³⁴] hPTH(1-34)NH₂ and [D-Bpa⁸, Tyr³⁴] hPTH(1-34)NH₂.

In yet another aspect, this invention provides a compound of formula (V),

 $(R^1R^2) - A^1 - A^2 - A^3 - A^4 - A^5 - A^6 - A^7 - A^8 - A^9 - A^{10} - A^{11} - A^{12} - A^{13} - A^{14} - A^{15} - A^{16} - A^{17} - A^{18} -$

20 $A^{19} - A^{20} - A^{21} - A^{22} - A^{23} - A^{24} - A^{25} - A^{26} - A^{27} - A^{28} - A^{29} - A^{30} - A^{31} - A^{32} - A^{33} - A^{34} - A^{35} - A^{36} - A^{37} - A^{38} - R^{3}$,

(V)

or a pharmaceutically acceptable salt thereof, wherein A^1 is Ala, Ser, Dap, Thr, Aib or is deleted;

25 A² is Val or is deleted;

A³ is Ser, Aib, Thr or is deleted;

A4 is Glu, Asp or is deleted;

 A^5 is His, Ile, Acc, Val, Nle, Phe, Leu, p-X-Phe, β -Nal, Aib, Cha or is deleted;

- 30 A⁶ is Gln, a hydrophilic amino acid or is deleted; A⁷ is Leu, Val, Cha, Nle, β -Nal, Trp, Pal, Acc, Phe, p-X-Phe, Aib, a lipophilic amino acid or is deleted; A⁸ is Leu, Met, Acc, Cha, Aib, Nle, Phe, Ile, Val, β -Nal, p-X-Phe, a lipophilic amino acid or is deleted;
- 35 A⁹ is His, a hydrophilic amino acid or is deleted; A¹⁰ is Asp, Asn, a hydrophilic amino acid or is deleted;

A¹¹ is Lys, Arg, Leu, Cha, Aib, p-X-Phe, Ile, Val, Nle, Acc, Phe, β -Nal, HN-CH((CH₂)_nNH-R⁴)-C(O), a lipophilic D-amino acid, a hydrophilic amino acid or is deleted;

A¹² is Gly, Acc, Aib or is deleted;

5 A^{13} is Lys, Arg, HN-CH((CH₂)_rNH-R⁴)-C(O) or is deleted; A^{14} is Ser, His or is deleted; A^{15} is Ile, Acc, Cha, Leu, Phe, Nle, β -Nal, Trp, p-X-Phe, Val, Aib or is deleted;

A¹⁶ is Gln, Aib or is deleted;

10 A^{17} is Asp, Aib or is deleted; A^{13} is Leu, Aib, Acc, Cha, Phe, Ile, Nle, β -Nal, Val, p-X-Phe or is deleted;

 A^{19} is Arg, Lys, Aib, $HN-CH((CH_2)_nNH-R^4)-C(0)$ or is deleted; A^{20} is Arg, Lys, $HN-CH((CH_2)_nNH-R^4)-C(0)$ or is deleted;

15 A^{21} is Arg, Lys, $HN-CH((CH_2)_nNH-R^4)-C(0)$ or is deleted; A^{22} is Phe, Glu, Aib, Acc, p-X-Phe, β -Nal, Val, Leu, Ile, Nle or Cha;

 A^{23} is Phe, Leu, Lys, Acc, Cha, β -Nal, Aib, Nle, Ile, p-X-Phe, Val or Trp;

20 A^{24} is Leu, Lys, Acc, Nle, Ile, Val, Phe, β -Nal, Aib, p-X-Phe, Arg or Cha;

A²⁵ is His, Lys, Aib, Acc, Arg or Glu;

A²⁶ is His, Aib, Acc, Arg or Lys;

 A^{27} is Leu, Lys, Acc, Arg, Ile, Val, Phe, Aib, Nle, β -Nal,

25 p-X-Phe or Cha;

 A^{28} is Ile, Leu, Lys, Acc, Cha, Val, Phe, p-X-Phe, Nle, β -Nal, Aib or is deleted;

A²⁹ is Ala, Glu, Acc, Aib or is deleted;

A³⁰ is Glu, Leu, Nle, Cha, Aib, Acc, Lys, Arg or is deleted;

30 A^{31} is Ile, Leu, Cha, Lys, Acc, Phe, Val, Nle, β -Nal, Arg or is deleted;

A32 is His or is deleted;

A³³ is Thr, Ser or is deleted;

 A^{34} is Ala, Phe, Tyr, Cha, Val, Ile, Leu, Nle, β -Nal, Aib,

35 Acc or is deleted;

A³⁵ is Glu, Asp or is deleted;

 A^{36} is Ile, Acc, Cha, Leu, Phe, Nle, β -Nal, Trp, p-X-Phe, Val, Aib or is deleted;

 A^{37} is Arg, Lys, $HN-CH((CH_2)_nNH-R^4)-C(0)$ or is deleted; A^{38} is Ala, Phe, Tyr, Cha, Val, Ile, Leu, Nle, β -Nal, Aib, Acc or is deleted;

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R¹ and R² are each independently selected from the group consisting of H, (C_{1-30}) alkyl, (C_{2-30}) alkenyl, phenyl- (C_{1-30}) alkyl, naphthyl (C_{1-30}) alkyl, hydroxy (C_{1-30}) alkyl, hydroxy (C_{2-30}) alkenyl, hydroxy-phenyl (C_{1-30}) alkyl or hydroxy-naphthyl (C_{1-30}) alkyl; or one of R¹ or R² is COE^1 where E^1 is (C_{1-30}) alkyl, (C_{2-30}) alkenyl, phenyl (C_{1-30}) alkyl, naphthyl (C_{1-30}) alkyl, hydroxy (C_{1-30}) alkyl, hydroxy-phenyl (C_{1-30}) alkyl, hydroxy-naphthyl (C_{1-30}) alkyl;

 R^3 is OH, NH_2 , (C_{1-30}) alkoxy or $NH-Y-CH_2-Z$, where Y is a (C_{1-30}) hydrocarbon moiety and Z is CO_2H or $CONH_2$; n for each occurrence is independently an integer from 1 to 5; and

 R^4 for each occurrence is independently (C_1 - C_{30}) alkyl, (C_1 - C_{30}) acyl or -C((NH)(NH₂));

provided that when A⁸ is not a lipophilic D-amino acid or is not deleted then at least one of A⁶, A⁷, A⁹, A¹⁰, A¹¹ and A¹² is a D-amino acid or at least one of A⁶, A⁷, A⁹, A¹⁰, A¹¹, A¹², A¹³, A¹⁴, A¹⁵, A¹⁶, A¹⁷, A¹⁸, A¹⁹, A²⁰, A²¹ and A²² is deleted.

A preferred group of compounds of formula (V) are the compounds listed as Examples 74-86, shown hereinbelow.

In a further aspect, this invention provides a method of selectively binding the PTH2 receptor which comprises administering to a patient in need thereof an analogue of formula (I), (II) or (III) or a pharmaceutically acceptable salt thereof.

In another aspect, this invention provides a method of selectively binding the PTH2 receptor which comprises administering to a patient in need thereof a compound of formula (III) or (V) or a pharmaceutically acceptable salt thereof. Preferred of the foregoing method is where the compound is selected from Examples 1-73 or Examples 74-86.

In another aspect, this invention is directed to a pharmaceutical composition comprising an analogue of

formula (I), (II) or (III) or a pharmaceutically acceptable salt thereof and a pharmaceutically acceptable carrier.

In still another aspect, this invention is directed to a pharmaceutical composition comprising a compound of formula (III) or (V) or a pharmaceutically acceptable salt thereof and a pharmaceutically acceptable carrier. Preferred is a pharmaceutical composition comprising a compound selected from Examples 1-73 or Examples 74-86.

In still another aspect, this invention is directed 10 to a method of treating a medical disorder that results from altered or excessive action of the PTH2 receptor, which comprises administering to a patient in need thereof an effective amount of a PTH analogue or a truncated PTH analogue or a pharmaceutically acceptable salt thereof that selectively binds to the PTH2 receptor, sufficient to inhibit the activation of the PTH2 receptor of said patient. A preferred method of the immediately foregoing method is where said medical disorder is abnormal CNS functions, abnormal pancreatic functions, divergence from metabolism and homeostasis, 20 normal mineral infertility, abnormal blood pressure or a hypothalmic disease. Preferred of each of the immediately foregoing methods is where the analogue is a PTH2 agonist or a PTH2 antagonist.

In another aspect, this invention provides a method of treating a medical disorder that results from altered or excessive action of the PTH2 receptor, which comprises administering to a patient in need thereof an effective amount of an analogue of formula (I), (II) or (III), sufficient to inhibit the activation of the PTH2 receptor of said patient. A preferred method of the immediately foregoing method is where said medical disorder is abnormal CNS functions, abnormal pancreatic functions, divergence from normal mineral metabolism and homeostasis, male infertility, abnormal blood pressure or a hypothalmic disease.

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In another aspect, this invention is directed to a method of treating a medical disorder that results from

altered or excessive action of the PTH2 receptor, which comprises administering to a patient in need thereof an effective amount of a compound of formula (III) or (V), sufficient to inhibit the activation of the PTH2 receptor of said patient. A preferred method of the immediately foregoing method is where said medical disorder is abnormal CNS functions, abnormal pancreatic functions, divergence from normal mineral metabolism and homeostasis, male infertility, abnormal blood pressure or a hypothalmic disease. Preferred of each of the foregoing methods is where the compound is selected from Examples 1-73 or Examples 74-86.

Detailed Description

With the exception of the N-terminal amino acid, all 15 abbreviations (e.g. Ala or A_1) of amino acids in this disclosure stand for the structure of -NH-CH(R)-CO-, wherein R is the side chain of an amino acid (e.g., CH, for Ala). For the N-terminal amino acid, the abbreviation stands for the structure of $(R^1R^2)-N-CH(R)-CO-$, wherein R is 20 a side chain of an amino acid and R1 and R2 are as defined above. Bpa is p-benzoylphenylalanine. β -Nal, Nle, Dap, Cha, Nva, Amp, Pal, and Aib are the abbreviations of the following α -amino acids: β -(2-naphthyl)alanine, norleucine, α, β -diaminopropionic acid, cyclohexylalanine, norvaline, 4-25 amino-phenylalanine, β -(3-pyridinyl)alanine aminoisobutyric acid, respectively. What is meant by Acc is an amino acid selected from the group of 1-amino-1cyclopropanecarboxylic acid; 1-amino-1cyclobutanecarboxylic acid; 1-amino-1-1-amino-1-30 cyclopentanecarboxylic acid; cyclohexanecarboxylic 1-amino-1acid; cycloheptanecarboxylic acid; 1-amino-1cyclooctanecarboxylic acid; and 1-amino-1cyclononanecarboxylic acid. In the above formula, 35 hydroxyalkyl, hydroxyphenylalkyl, and hydroxynaphthylalkyl may contain 1-4 hydroxy substituents. COE, stands for - $C=0\cdot E^1$. Examples of $-C=0\cdot E^1$ include, but are not limited to, acetyl and phenylpropionyl. What is meant by " (C_{1-12})

hydrocarbon moiety" is an alkyl group, an alkenyl group or an alkynyl group.

What is meant by a "hydrophilic amino acid" is an amino acid having at least one hydrophilic functional group in addition to those required for peptide bond formation, such as: Arg, Asp, Asn, Glu, Gln, Gly, His, Lys, Orn (ornithine), Ser, Thr, β -Ala, Ala, Aad (α -aminoadipic acid), β -Aad (β -aminoadipic acid), Apm (α -aminopimolic acid), Cit (citrulline), Gla (γ -carboxy-glutamic acid), hArg (homo-Arg), hCit (homo-Cit), hSer (homo-Ser), Dba (α , γ -diamino-butyric acid), Dpa (α , β -diaminopropionic acid), Amp (β -amino-phenylalanine), Pal, and their homologues.

What is meant by a "lipophilic amino acid" is an uncharged, aliphatic or aromatic amino acid, such as: Val, Leu, Ile, Pro, Cys, Phe, Met, Trp, Tyr, Cha, β-Nal, Aib, Acc, Ala, Abu (α-aminobutyric acid), Nle, Nva (norvaline), Bpa (p-benzoyl-phenylalanine), hPhe (homo-Phe), hPro (homo-Pro), 1-Nal(β-(1-naphthyl)alanine), 2-Nal (β(2-naphthyl)alanine), Oic (octahydroindode-2-carboxylic acid), Tic (1,2,3,4-tetrahydroisoquinoline-3-carboxylic acid), Pen (penicillamine), Phg (phenylglycine), Tle (t-leucine), p-X-Phe (X= Br, F, I, Cl, CH, phenyl, CN, NO₂), Tal (β-(2-thienyl)-alanine), and their homologues.

Alanine, β -alanine and sarcosine (Sar) may be considered either a hydrophilic or a lipophilic amino acid.

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"Physiologically active truncated homologue or analogue of PTH" refers to a polypeptide having a sequence comprising less than the full complement of amino acids found in PTH.

The full names for other abbreviations used herein are as follows: Boc for t-butyloxycarbonyl, HF for hydrogen fluoride, Fm for formyl, Xan for xanthyl, Bzl for benzyl, Tos for tosyl, DNP for 2,4-dinitrophenyl, DMF for dimethylformamide, DCM for dichloromethane, HBTU for 2-(1H-Benzotriazol-1-yl)-1,1,3,3-tetramethyl uronium hexafluorophosphate, DIEA for diisopropylethylamine, HOAc

for acetic acid, TFA for trifluoroacetic acid, 2ClZ for 2chlorobenzyloxycarbonyl and OcHex for O-cyclohexyl.

A peptide of this invention is also denoted herein by another format, e.g., [D-Nle8] hPTH(1-34)NH2, with the 5 substituted amino acids from the natural sequence placed between the set of brackets (e.g., D-Nle3 for Met8 in hPTH). The abbreviation hPTH stands for human PTH, and hPTHrP for human PTHrP. The numbers between the parentheses refer to the number of amino acids present in the peptide (e.g., 10 hPTH(1-34) is amino acids 1 through 34 of the peptide sequence for human PTH). The sequences for hPTH(1-34) and hpTHrP(1-34) are listed in Nissenson, et al., Receptor, The designation " NH_2 " in $PTH(1-34)NH_2$ 3:193 (1993). indicates that the C-terminus of the peptide is amidated. PTH(1-34) means that the C-terminus is the free acid.

The peptides of this invention can be prepared by standard solid phase peptide synthesis. See, e.q., Stewart, J.M., et al., Solid Phase Synthesis (Pierce Chemical Co., 2d ed. 1984). The substituents R^1 and R^2 of 20 the above generic formula may be attached to the free amine of the N-terminal amino acid by standard methods known in the art. For example, alkyl groups, e.g., (C_{1-12}) alkyl, may be attached using reductive alkylation. Hydroxyalkyl groups, e.g., (C_{1-12}) hydroxyalkyl, may also be attached using reductive alkylation wherein the free hydroxy group is protected with a t-butyl ester. Acyl groups, e.g., COE1, may be attached by coupling the free acid, e.g., E1COOH, to the free amine of the N-terminal amino acid by mixing the completed resin with 3 molar equivalents of both the free acid and diisopropylcarbodiimide in methylene chloride for one hour. If the free acid contains a free hydroxy group, e.g., p-hydroxyphenylpropionic acid, then the coupling should be performed with an additional 3 molar equivalents of HOBT.

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When R³ is NH-Y-CH₂-CONH₂ (Z=CONH₂), the synthesis of 35 the peptide starts with BocHN-Y-CH2-COOH which is coupled to the resin. If R^3 is NH-Y-CH₂-COOH (Z=COOH) the synthesis of the peptide starts with $Boc-HN-Y-CH_2-COOH$ which is coupled to PAM resin. When R^3 is OH the first amino acid is coupled to PAM resin.

The compounds of this invention can be tested for binding to the human PTH2 (hPTH2) receptor for the ability to stimulate adenylyl cyclase and/or intracellular calcium transients by the assay described below.

Materials and Methods: Tissue culture media and sera were purchased from Life Technologies (Grand Island, NY), and all tissue culture plastics were obtained from 10 Corning (Corning, NY). Adenosine and 3-isobutyl-1-methyl xanthine (IBMX) were purchased from Research Biochemicals (Natick, MA). Fura-2 acetoxylmethyl ester (fura-2/AM) was obtained from Molecular Probes (Eugene, OR), and hPTHrP was purchased from Bachem (Torrance, CA). [3H]-Adenine was purchased from New England Nuclear (Boston, MA). Na¹²⁵I was obtained from Amersham Corp. (Arlington Heights, IL). All other analytical grade reagents were purchased from Sigma (St. Louis, MO).

Cell Culture: Human osteosarcoma Saos-2/B-10 cells 20 (American Type Culture Collection, Rockville, MD; ATCC #HTB 85) are maintained in RPMI 1640 medium (Sigma, St. Louis, MO) supplemented with 10% fetal bovine serum (FBS) and 2 mM glutamine at 37°C in a humidified atmosphere of 5% CO2 in air. The medium is changed every three or four days, and 25 the cells are subcultured every week by trypsinization. Stably transfected HEK-293/BP-16 cells (Beth Deaconess Medical Center-Division of Bone and Mineral Metabolism, Boston, MA), which express the hPTH2 receptor (160,000 receptors/cell) and stably transfected HEK-293/C-21 cells (Beth Israel Deaconess Medical Center-Division of Bone and Mineral Metabolism, Boston, MA), which express the hPTH/PTHrP receptor, are maintained in DMEM supplemented with 10% FBS at 37°C in a humidified atmosphere of 95% air/5% CO2. The medium is changed every 2 days before confluency and every day after confluency. The cells sub-cultured 1:10 once a week.

Receptor binding assay: Ligand binding is performed using Saos-2/B-10, HEK/C-21 cells or HEK/BP-16 cells using

HPLC-purified [125I] [$Nle^{8.18}$, Tyr^{34}] bPTH-(1-34) NH_2 (125I-PTH) as radioligand. Saos-2 cells are maintained for four days until they reach confluence. The medium is replaced with 5% FBS in RPMI 1640 medium and incubated for about 2 hrs at 5 room temperature with 10 x 10 4 cpm mono- ^{125}I -[Nle 8,18 , Tyr 34 (3-125I)]bPTH(1-34)NH, in the presence of competing peptides of the invention at various concentrations between 10⁻¹¹M to 10⁻ The cells are washed four times with ice-cold PBS and lysed with 0.1 M NaOH, and the radioactivity associated with the cells is counted in a scintillation counter. Synthesis of mono- ^{125}I -[Nle 8,18 , Tyr 34 (3- ^{125}I)]bPTH(1-34)NH, is carried out as described in Goldman, M.E., et al., Endocrinol., 123:1468 (1988).

The binding assay is conducted with various peptides of the invention, and the Kd value (half maximal inhibition of binding of mono- ^{125}I -[Nle 8,18 , Tyr 34 (3- ^{125}I)]bPTH(1-34)NH₂) for each peptide is calculated.

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Adenylyl cyclase assay: Adenylyl cyclase assay is performed in Saos-2/B-10 cells, HEK/C21 cells, and HEK/BP-16 cells. The ability of the peptides of the invention to induce a biological response in Saos-2/B-10 cells is More specifically, any stimulation of the adenylate cyclase is determined by measuring the level of synthesis of cAMP (adenosine 3',5'-monophosphate) described previously in Rodan, et al., J. Clin. Invest. 72: 1511 (1983) and Goldman, et al., Endocrinol., 123:1468 (1988). Confluent Saos-2/B-10 cells in 24 well plates at 4x104 cells/well in RPMI1640 medium containing 10% FBS. Cells are washed twice with Ca2+ and Mg2+ free Hanks' 30 balanced salt solution and incubated with [3H] adenine (26.9 Ci/mmol, New England Nuclear, Boston, MA) in fresh medium at about 37°C for about 2 hrs, and washed Hank's balanced salt solution (Gibco, twice with Gaithersburg, MD). The cells are treated with 1 mM IBMX [isobutylmethyl-xanthine, Sigma, St. Louis, MO] in fresh medium for 15 min, and a peptide to be tested is added to the medium to incubate for about 5 min. The reaction is stopped by the addition of 1.2 M trichloroacetic acid (TCA)

(Sigma, St. Louis, MO) followed by sample neutralization cAMP is isolated by the two-column with 4 N KOH. et al., 1974, chromatographic method (Salmon, Biochem. 58, 541). The radioactivity is counted in a 5 scintillation counter (Liquid Scintillation Counter 2200CA, PACKARD, Downers Grove, IL).

Measurements [Ca²⁺],: Measurements of of intracellular Ca2+ ([Ca2+]) are performed in Saos-2/B-10 cells, HEK/C-21 cells and HEK/BP-16 cells. For measurement of [Ca²⁺], cells are harvested from 150-cm² flasks using HEPES-buffered balanced salt solution containing 0.02% (vol/vol) EDTA. The cell suspension is washed three times with Hanks' Balanced Salt Solution (1 mM CaCl2, 118 mM NaCl, 4.6 mM KCl, 10 mM d-glucose, and 20 mM HEPES, pH 7.4), and 15 cells are loaded with fura-2/AM (1 μ M) for about 40 min at about 37°C. The cell suspension is washed three times with Hanks' Balanced Salt Solution, and fluorescence is measured in a SPEX AR-CM system spectrofluorimeter (SPEX Industries, Edison, NJ). Dual wavelength measurements are performed (excitation wavelengths, 340 and 380 nm; emission wavelength, 505 nm).

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[Ca²⁺]_i is calculated from fura-2 ratios (R) by the $[Ca^{2+}]_i = K (R - R_{min}) / (R_{max} - R)$, where R_{min} and R_{max} are the ratios (e.g. 340 nm/380 nm) for the minimal or maximal calcium concentration, respectively. product $K_d(F_o/F_s)$, where K_d is the effective dissociation constant (224 nM), F_0 is the intensity of the 380-nm excitation signal in the absence of calcium, and $F_{\rm s}$ is the intensity of the 380-nm excitation signal at saturating calcium concentrations. Maximum fluorescence intensity is obtained by permeabilizing the cells with 50 μM digitonin in the presence of 1 mM CaCl2, and minimal fluorescence intensity is obtained by chelating calcium with 16.6 mM 8.3 with 1M to **EGTA** Hq] adjusted (hydroxymethyl)aminomethane base]. Addition of vehicle alone (0.1% BSA in PBS) did not change the level of [Ca2+];.

The peptides of this invention can be provided in the form of pharmaceutically acceptable salts. Examples of

such salts include, but are not limited to, those formed with organic acids (e.g., acetic, lactic, maleic, citric, malic, ascorbic, succinic, benzoic, methanesulfonic, toluenesulfonic or pamoic acid), inorganic acids (e.g., hydrochloric acid, sulfuric acid, or phosphoric acid), and polymeric acids (e.g., tannic acid, carboxymethyl cellulose, polylactic, polyglycolic, or copolymers of polylactic-glycolic acids).

A therapeutically effective amount of a peptide of 10 this invention and a pharmaceutically acceptable carrier substance (e.g., magnesium carbonate, lactose, phospholipid with which the therapeutic compound can form a micelle) together form a therapeutic composition (e.g., a pill, tablet, capsule, or liquid) for administration (e.g., orally, intravenously, transdermally, pulmonarily, 15 vaginally, subcutaneously, nasally, iontophoretically, or by intratracheally) to a subject. The pill, tablet or capsule that is to be administered orally can be coated with a substance for protecting the active composition from 20 the gastric acid or intestinal enzymes in the stomach for a period of time sufficient to allow it to pass undigested into the small intestine. The therapeutic composition can also be in the form of a biodegradable or nonbiodegradable release formulation for subcutaneous sustained See, e.g., U.S. Patents intramuscular administration. 25 3,773,919 and 4,767,628 and PCT Application No. Continuous administration can also be achieved 94/15587. using an implantable or external pump (e.g., INFUSAID™ administration can also be conducted pump). The 30 intermittently, e.g., single daily injection, continuously at a low dose, e.g., sustained release formulation.

Liquid dosage forms for oral administration include pharmaceutically acceptable emulsions, solutions, suspensions, syrups, the elixirs containing inert diluents commonly used in the art, such as water. Besides such inert diluents, compositions can also include adjuvants,

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such as wetting agents, emulsifying and suspending agents, and sweetening, flavoring and perfuming agents.

Preparations according to this invention parenteral administration include sterile aqueous or non-5 aqueous solutions, suspensions, or emulsions. Examples of non-aqueous solvents or vehicles are propylene glycol, polyethylene glycol, vegetable oils, such as olive oil and corn oil, gelatin, and injectable organic esters such as ethyl oleate. Such dosage forms may also contain adjuvants such as preserving, wetting, emulsifying, and dispersing agents. They may be sterilized by, for example, filtration through a bacteria-retaining filter, by incorporating sterilizing agents into the compositions, by irradiating the compositions, or by heating the compositions. also be manufactured in the form of sterile solid compositions which can be dissolved in sterile water, or some other sterile injectable medium immediately before use.

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Compositions for rectal or vaginal administration are preferably suppositories which may contain, in addition to the active substance, excipients such as coca butter or a suppository wax.

Compositions for nasal or sublingual administration are also prepared with standard excipients well known in the art.

Further, a compound of this invention can be administered in a sustained release composition such as those described in the following patents. U.S. Patent No. 5,672,659 teaches sustained release compositions comprising a bioactive agent and a polyester. U.S. Patent 5,595,760 teaches sustained release compositions comprising a bioactive agent in a gelable form. U.S. Application No. 08/929,363 filed September 9, 1997, teaches polymeric sustained release compositions comprising a bioactive agent chitosan. U.S. Application No. 08/740,778 November 1, 1996, teaches sustained release compositions comprising a bioactive agent and cyclodextrin. U.S. Application No. 09/015,394 filed January 29, 1998, teaches

absorbable sustained release compositions of a bioactive The teachings of the foregoing patents applications are incorporated herein by reference.

The dosage of active ingredient in the compositions 5 of this invention may be varied; however, it is necessary that the amount of the active ingredient be such that a suitable dosage form is obtained. The selected dosage depends upon the desired therapeutic effect, on the route of administration, and on the duration of the treatment. 10 Generally, dosage levels of between 0.0001 to 10 mg/kg of body weight daily are administered.

A preferred dosage range is 0.001 to 0.5 mg/kg of body weight daily which can be administered as a single dose or divided into multiple doses.

15 The compounds of the instant invention illustrated by the following examples, but are not limited to the details thereof.

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EXAMPLE 1

[Cha 7,11 , D-Nle 8 , Nle 18 , Tyr 34]hPTH(1-34)NH₂

The peptide [Cha^{7,11}, D-Nle⁸, Nle¹⁸, Tyr³⁴] hPTH (1-34) NH, was synthesized on an Applied Biosystems (Foster City, CA) model 430A peptide synthesizer which was modified to do accelerated Boc-chemistry solid phase peptide synthesis. See Schnoize, et al., Int. J. Peptide Protein Res., 90:180 (1992). 4-Methylbenzhydrylamine (MBHA) resin (Peninsula, Belmont, CA) with the substitution of 0.93 mmol/g was used. (Bachem, CA, Torrance, The Boc amino acids Biochem., LaJolla, CA) were used with the following side chain protection: Boc-Asn(Xanthyl), Boc-Arg(Tos)-OH, Boc-30 Asp(OcHex)-OH, Boc-Glu(OcHex)-OH, Boc-His(DNP)-OH, Boc-Cha-OH, Boc-D-Nle-OH, Boc-Nle-OH, Boc-Val-OH, Boc-Leu-OH, Boc-Gly-OH, Boc-Gln-OH, Boc-Ile-OH, Boc-Lys(2ClZ)-OH, Ser(Bzl)-OH; Boc-Trp(formyl)-OH and Boc-Tyr(Br-Z)-OH (where Z is benzyloxycarbonyl). The synthesis was carried out on a 0.14 mmol scale. The Boc groups were removed by treatment with 100% TFA for 2 x 1 min. Boc amino acids (2.5 mmol) were pre-activated with HBTU (2.0 mmol) and DIEA (1.0 mL) 4 mLof DMF and were coupled without in

neutralization of the peptide-resin TFA salt. Coupling times were about 5 min.

At the end of the assembly of the peptide chain, the treated with а solution 20% was resin mercaptoethanol/10% DIEA in DMF for 2 x 30 min. to remove the DNP group on the His side chain. The resin was washed with DMF. The N-terminal Boc group was then removed by treatment with 100% TFA for 2 x 2 min. The resin was washed with DMF and was treated with ethanolamine: H2O:DMF/15:15:70 10 for 2 \times 30 min. to remove the formyl protecting group on Trp residue. The partially-deprotected peptide-resin was washed with DMF and DCM and dried in vacuo. The final cleavage was done by stirring the peptide-resin in 10 mL of HF containing 1 mL of anisole and dithiothreitol (24 mg) at 15 about 0°C for about 75 min. HF was removed by a flow of nitrogen. The residue was washed with ether (6 x 10 mL) and extracted with 4N HOAc (6 x 10 mL).

The peptide mixture in the aqueous extract was purified on a reverse-phase preparative high pressure liquid chromatography (HPLC) using a reverse phase VYDACTM C₁₈ column (Nest Group, Southborough, MA). The column was eluted with a linear gradient (10% to 45% of solution B in solution A over 130 min.) at a flow rate of 10 mL/min (Solution A = water containing 0.1% TFA; Solution B = acetonitrile containing 0.1% of TFA). Fractions were collected and checked on analytical HPLC. Those containing pure product were combined and lyophilized to dryness. 114 mg of a white solid was obtained. Purity was >98% based on analytical HPLC analysis. Electro-spray mass spectrometer analysis gave the molecular weight at 4176.4 (in agreement with the calculated molecular weight of 4176.9).

EXAMPLE 2

[D-Nle⁸, Nle¹⁸, Tyr³⁴] hPTH (1-34) NH₂

Boc-protected amino acids, N-hydroxybenzotriazole 35 (HOBt), N,N'-dicyclohexylcarbodilmide (DCC) and p-methylbenzhydrylamine resin were purchased from Applied Biosystems (Foster City, CA). Boc-(3-Iodo)Tyrosine[0-(3-BrBz)] was purchased from Peninsula Laboratories (Belmont,

CA). B&J brand dichloromethane, N-methylpyrrolidone (NMP) and acetonitrile were obtained from Baxter (McGraw Park, IL). All other reagents are commercially available, for example from Sigma (St. Louis, MO). The title peptide was 5 synthesized by solid-phase Boc/HOBt/NMP chemistry on an automated Applied Biosystems 430A peptide synthesizer using software version 1.40. The following side-chain protected $N-\alpha$ -Boc-amino derivatives were used in the course of the automated solid-phase peptide synthesis: Arg(No-tosyl), Asp(O-cHex), Glu(O-Bzl), His(Nn-Bom), Lys(N'-2-Cl-Z), Ser(O-10 Bzl), Thr(O-Bzl), and Tyr(2-Br-Z). Synthesis started at a 0.5 mmol scale and was split into two halves after the incorporation of Glu²². The following residues were incorporated by double coupling cycles: Arg25, Leu24, Val21, Arg²⁰, Glu¹⁹, Leu¹⁵, His¹⁴, Lys¹³, His⁹, Phe⁷, Gln⁶ and Ile⁵. The Nle in positions 18 and 8 was introduced in the form of pre-dissolved NMP solution and the Activator cycle was modified accordingly. Cleavage of the peptide from the $ho \mathrm{MBHA}$ resin utilized liquid hydrogen fluoride and followed The "Low-HF" step included the "Low-High" procedure. 20 mixing the suspension of the resin-bound peptide in a mixture (20 mL/q of resin-bound peptide) containing (% vol) 60% dimethylsulfide, 5% ρ -thiocresol, 5% ρ -cresol, ethane dithiol, and 25% HF for about 2 hours at about 0°C. After removal of the volatile reagent under vacuum and the resin-bound peptide consecutively with petroleum-ether and ether it was returned to the reaction vessel for the "High-HF" step. The resin-bound peptide was resuspended in a mixture (20 mL/g of resin-bound peptide) 30 containing (% vol) 5% butane dithiol, 5% ρ -cresol, and 90% HF for about 1 hour at about O°C. After removing the reagents as previously described the crude peptide was dissolved in 50% (v/v) acetic acid and the solution was diluted with water and lyophilized. The peptide purified by preparative reverse-phase high performance liquid chromatography (RP-HPLC) (PrepPak VYDAC® C18, 300Å cartridge, 15 μ m, 5.5x35 cm). The solvent system employed included a two solvent system: A: 0.1% (v/v) TFA in water and B: 0.1% (v/v) TFA in acetonitrile, generating

the following linear gradient: 0-15% B in A in the first 10 min followed by 15-45% B in A in the next 120 min at a flow-rate of 70 mL/min and monitored at 220 nm. Fractions were analyzed on an analytical RP-HPLC system (VYDAC® (C18, 300Å, 5 μ m, 4.6x150cm) employing a linear gradient of 20-50% B in A for 30 min at a flow rate of 1 ml/min and monitored at 220 nm, the retention time is 18.24 minutes. The pure fractions were pooled and the acetonitrile removed under vacuum. The residual was lyophilized to yield a white powder. Purity and structure of the peptides were confirmed by analytical RP-HPLC, amino acid analysis, and Fast Atom Bombardment Mass Spectrometry, mass spec. = 4097.0.

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EXAMPLES 3-5

Examples 3-4 were synthesized substantially according to the procedure of Example 1 using the appropriate, protected amino acids and Example 5 was synthesized substantially according to Example 2 using the appropriate, protected amino acids.

15	Example	Name	Mass Spec.
	3	[Cha ^{7,17} , des-Met ⁸ , Nle ¹⁸ , Tyr ³⁸]hPTH(1-34)NH ₂	4063.5
	4	[Cha'.'', D-Nle*, des-Met'*, Tyr**]hPTH(1-34)NH ₂	4063.4
	5	[D-Bpa ^s , Tyr ^{ss}]hPTH-(1-34)NH ₂	4320.7

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EXAMPLES 6-86

Examples 6 to 86 can be synthesized substantially according to the procedure of Example 1 using the appropriate, protected amino acids.

Example 6:

[D-Nie⁸, Nie¹⁸, Tyr³⁴]hPTH(1-34)NH₂

Example 7:

[D-Nle⁸]hPTH(1-34)NH₂

25 Example 8:

[D-Leu⁸, Nie¹⁸, Tyr³⁴]hPTH(1-34)NH₂

Example 9:

[D-Cha⁸, Nie¹⁸, Tyr³⁴]hPTH(1-34)NH₂

Example 10:

[D-Phe⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂

Example 11:

[D-Nal⁸, Nie¹⁸, Tyr³⁴]hPTH(1-34)NH₂

Example 12:

[D-Abu⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂

30 Example 13:

[D-Met⁸]hPTH(1-34)NH₂

Example 14:

[Cha^{7, 11}, D-Met⁸]hPTH(1-34)NH₂

Example 15:

[D-lle⁸]hPTH(1-34)NH₂

Example 16:

[Cha^{7, 11}, D-lle⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂

Example 17:

[D-IIe8, NIe18, Tyr34]hPTH(1-34)NH,

35 Example 18:

[D-Leu⁸]hPTH(1-34)NH₂

WO 99/57139 PCT/US99/09521 30

[Cha^{7,11}, D-Leu⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂ Example 19: Example 20: [D-Val⁸]hPTH(1-34)NH₂ [Cha^{7, 11}, D-Val⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂ Example 21: [D-Val⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂ Example 22: [D-Cha⁸]hPTH(1-34)NH₂ 5 Example 23: [Cha^{7,11}, D-Cha⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂ Example 24: [D-Ala8]hPTH(1-34)NH₂ Example 25: [Cha^{7, 11}, D-Ala⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂ Example 26: [D-Ala⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂ Example 27: Example 28: [D-Phe⁸]hPTH(1-34)NH₂ 10 [Cha^{7,11}, D-Phe⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂ Example 29: [D-Met⁸]hPTH(7-34)NH₂ Example 30: [D-Nal8]hPTH(1-34)NH₂ Example 31: [D-Trp⁸]hPTH(1-34)NH₂ Example 32: [Cha^{7,11}, D-Trp⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂ Example 33: 15 [D-Trp8, Nle18, Tyr34]hPTH(1-34)NH₂ Example 34: [D-Abu8]hPTH(1-34)NH₂ Example 35: [Cha^{7,11}, D-Abu⁸, Nie¹⁸, Tyr³⁴]hPTH(1-34)NH₂ Example 36: [des-Met⁸]hPTH(1-34)NH₂ Example 37: [Cha^{7,11}, des-Met⁸]hPTH(1-34)NH₂ 20 Example 38: [Cha^{7,11}, des-Met⁸, des-Met¹⁸, Tyr³⁴]hPTH(1-34)NH₂ Example 39: [des-Met⁸, des-Met¹⁸]hPTH(1-34)NH₂ Example 40: [Cha^{7,11}, des-Met⁸, des-Met¹⁸]hPTH(1-34)NH₂ Example 41: [des-Met⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂ Example 42: [des-Met¹⁸]hPTH(1-34)NH₂ Example 43: 25 [Cha^{7,11}, des-Met¹⁸]hPTH(1-34)NH₂ Example 44: [Cha^{7,11}, des-Met¹⁸, Tyr³⁴]hPTH(1-34)NH₂ Example 45: [D-Nie⁸, des-Met¹⁸, Tyr³⁴]hPTH(1-34)NH₂ Example 46: [des-Glu⁶, Nle^{8,18}, Tyr³⁴]hPTH(1-34)NH₂ Example 47: [des-Leu⁷, Nle^{8,18}, Tyr³⁴]hPTH(1-34)NH₂ Example 48: 30 [des-His⁹, Nle^{8,18}, Tyr³⁴]hPTH(1-34)NH₂ Example 49: [des-Asn¹⁰, Nle^{8,18}, Tyr³⁴]hPTH(1-34)NH₂ Example 50: [des-Leu¹¹, Nle^{8,18}, Tyr³⁴]hPTH(1-34)NH₂ Example 51: [des-Giy¹², Nle^{8,18}, Tyr³⁴]hPTH(1-34)NH₂ Example 52: Ides-Lys¹³, Nle^{8,18}, Tyr³⁴lhPTH(1-34)NH₂ Example 53: 35

	Example 54:	[des-His 14 , Nie $^{8.18}$, Tyr 24]hPTH(1-34)NH $_2$
	Example 55:	[des-Leu ¹⁵ , Nle ^{8,18} , Tyr ³⁴]hPTH(1-34)NH ₂
	Example 56:	[des-Asn ¹⁶ , Nle ^{8,18} , Tyr ³⁴]hPTH(1-34)NH ₂
	Example 57:	[des-Ser 17 , Nle $^{8.18}$, Tyr 34]hPTH(1-34)NH $_2$
5	Example 58:	[des-Glu ¹⁹ , Nle ^{8,18} , Tyr ³⁴]hPTH(1-34)NH ₂
	Example 59:	[des-Arg ²⁰ , Nle ^{8,18} , Tyr ³⁴]hPTH(1-34)NH₂
	Example 60:	[des-Val 21 , Nle $^{8.18}$, Tyr 34]hPTH(1-34)NH $_2$
	Example 61:	[des-Glu 22 , Nle 8,18 , Tyr 34]hPTH(1-34)NH $_2$
	Example 62:	[des-Glu ⁶ , Cha ^{7,11} , Nle ^{8,18} , Tyr ³⁴]hPTH(1-34)NH ₂
10	Example 63:	[des-Leu ⁷ , Nle ^{8,18} , Cha ¹¹ , Tyr ³⁴]hPTH(1-34)NH ₂
	Example 64:	[Cha ^{7,11} , des-His ⁹ , Nle ^{8,18} , Tyr ³⁴]hPTH(1-34)NH ₂
	Example 65:	[des-Glu ⁶ , Cha ^{7,11} , D-Nle ⁸ , Nle ¹⁸ , Tyr ³⁴]hPTH(1-34)NH ₂
	Example 66:	[des-Leu ⁷ , D-Nle ⁸ , Cha ¹¹ , Nle ¹⁸ , Tyr ³⁴]hPTH(1-34)NH ₂
	Example 67:	[Cha ^{7,11} , D-Nle ⁸ , des-His ⁹ , Nle ¹⁸ , Tyr ³⁴]hPTH(1-34)NH ₂
15	Example 68:	[Cha ^{7,11} , des-Met ⁸ , des-His ⁹ , des-Asn ¹⁰]hPTH(1-34)NH ₂
	Example 69:	[Cha ^{7,11} , des-Ser ¹⁷ , des-Met ¹⁸ , des-Glu ¹⁹]hPTH(1-34)NH ₂
	Example 70:	[D-Met ⁸ , Nle ¹⁸ , Tyr ³⁴]hPTH(1-34)NH ₂
	Example 71:	[D-Met ⁸ , Tyr ³⁴]hPTH(1-34)NH₂
	Example 72:	[D-Nle 8 , Nle 18 , Tyr 34]hPTH(7-34)NH $_2$
20	Example 73:	[D-Nle ⁸ , Nle ¹⁸]hPTH(7-34)NH ₂
	Example 74:	[lle ⁵ , D-Leu ⁸]hPTHrP(1-34)NH ₂
	Example 75:	[lle ⁵ , D-Leu ⁸ , Trp ²³]hPTHrP(1-34)NH ₂
	Example 76:	[lle ⁵ , des-Leu ⁸ , Trp ²³]hPTHrP(1-34)NH ₂
	Example 77:	[lle ⁵ , des-Leu ⁸]hPTHrP(1-34)NH ₂
25	Example 78:	[des-Leu ⁸ , Trp ²³]hPTHrP(1-34)NH ₂
	Example 79:	[lle ⁵ , des-Leu ¹⁸]hPTHrP(1-34)NH ₂
	Example 80:	[lle ⁵ , des-Leu ¹⁸ , Trp ²³]hPTHrP(1-34)NH ₂
	Example 81:	[des-Leu ¹⁸ , Trp ²³]hPTHrP(1-34)NH ₂
	Example 82:	[lle ⁵ , D-Leu ⁸ , Glu ^{22,25} , Leu ^{23,28,31} , Lys ^{26,30} , Aib ²⁹]hPTHrP(1-34)NH ₂
30	Example 83:	[lle ⁵ , D-Leu ⁸ , Glu ^{22,25} , Trp ²³ , Lys ^{26,30} , Leu ^{28,31} , Aib ²⁹]hPTHrP(1-34)NH ₂
	Example 84:	[lle ⁵ , D-Leu ⁸ , Glu ^{22,25,29} , Leu ^{23,28,31} , Lys ^{26,30}]hPTHrP(1-34)NH ₂
	Example 85:	[lle ⁵ , D-Leu ⁸ , Glu ^{22,25,29} , Trp ²³ , Lys ^{26,30} , Leu ^{28,31}]hPTHrP(1-34)NH ₂
	Example 86:	[D-Leu ⁸ , Trp ²³]hPTHrP(7-34)NH ₂

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CLAIMS

What is claimed is:

- 1. A PTH analogue or a truncated PTH analogue or a pharmaceutically acceptable salt thereof that selectively binds to the PTH2 receptor.
- 2. A PTH analogue or a truncated PTH analogue or a pharmaceutically acceptable salt thereof according to claim 1 where said analogue is a selective PTH2 receptor agonist.
 - 3. A PTH analogue or a truncated PTH analogue or a pharmaceutically acceptable salt thereof according to claim 1 where said analogue is a selective PTH2 receptor antagonist.
 - 4. A method of selectively binding a PTH2 receptor which comprises administering to a patient in need thereof an effective amount of an analogue according to claim 1 or a pharmaceutically-acceptable salt thereof.
- 5. A method of selectively eliciting an agonist response from the PTH2 receptor which comprises administering to a patient in need thereof an effective amount of an analogue according to claim 2 or a pharmaceutically acceptable salt thereof.
 - 6. A method of selectively eliciting an antagonist response from the PTH2 receptor which comprises administering to a patient in need thereof an effective amount of an analogue according to claim 3 or a pharmaceutically acceptable salt thereof.
 - An analogue according to claim 1 wherein said analogue is of formula (I),

 $(\mathsf{R}^1\mathsf{R}^2) - \mathsf{A}^1 - \mathsf{A}^2 - \mathsf{A}^3 - \mathsf{A}^4 - \mathsf{A}^5 - \mathsf{A}^6 - \mathsf{A}^7 - \mathsf{A}^8 - \mathsf{A}^9 - \mathsf{A}^{10} - \mathsf{A}^{11} - \mathsf{A}^{12} - \mathsf{A}^{13} - \mathsf{A}^{15} - \mathsf{A}^{16} - \mathsf{A}^{17} - \mathsf{A}^{18} - \mathsf{A}^{19} - \mathsf{A}^{20} - \mathsf{A}^{21} - \mathsf{A}^{22} - \mathsf{A}^{23} - \mathsf{A}^{24} - \mathsf{A}^{25} - \mathsf{A}^{26} - \mathsf{A}^{27} - \mathsf{A}^{28} - \mathsf{A}^{29} - \mathsf{A}^{30} - \mathsf{A}^{31} - \mathsf{A}^{32} - \mathsf{A}^{33} - \mathsf{A}^{34} - \mathsf{A}^{35} - \mathsf{A}^{36} - \mathsf{A}^{37} - \mathsf{A}^{38} - \mathsf{R}^3$

(1)

or a pharmaceutically-acceptable salt thereof wherein

A¹ is a hydrophilic or a lipophilic amino acid;

A² is a lipophilic amino acid;

30 A³ is a hydrophilic or a lipophilic amino acid;

A⁴ is a hydrophilic amino acid;

A⁵ is a hydrophilic or a lipophilic amino acid;

A⁶ is a hydrophilic amino acid or is deleted;

A⁷ is a hydrophilic or a lipophilic amino acid or is deleted;

35 A⁸ is a lipophilic amino acid or is deleted;

A9 is a hydrophilic amino acid or is deleted;

A¹⁰ is a hydrophilic amino acid or is deleted;

A¹¹ is a hydrophilic or a lipophilic amino acid or is deleted;

A¹² is a hydrophilic or a lipophilic amino acid or is deleted;

5 A¹³ is a hydrophilic amino acid;

A¹⁴ is a hydrophilic amino acid or is deleted;

A¹⁵ is a lipophilic amino acid or is deleted;

A¹⁶ is a hydrophilic or a lipophilic amino acid or is deleted;

A¹⁷ is a hydrophilic or a lipophilic amino acid or is deleted;

10 A¹⁸ is a lipophilic amino acid or is deleted;

A¹⁹ is a hydrophilic or a lipophilic amino acid or is deleted;

A²⁰ is a hydrophilic amino acid or is deleted;

A²¹ is a hydrophilic or a lipophilic amino acid or is deleted;

A²² is a lipophilic or a hydrophilic amino acid or is deleted;

15 A²³ is a hydrophilic or a lipophilic amino acid;

A²⁴ is a hydrophilic or a lipophilic amino acid;

A²⁵ is a hydrophilic amino acid;

A²⁶ is a hydrophilic amino acid;

A²⁷ is a lipophilic or a hydrophilic amino acid;

20 A²⁸ is a lipophilic amino acid;

A²⁹ is a lipophilic or a hydrophilic amino acid;

A³⁰ is a hydrophilic or a lipophilic amino acid;

A³¹ is a lipophilic or a hydrophilic amino acid or is deleted;

A³² is a hydrophilic amino acid or is deleted;

25 A³³ is a hydrophilic amino acid or is deleted;

A³⁴ is a lipophilic amino acid or is deleted;

A³⁵ is a lipophilic amino acid or is deleted;

A³⁶ is a lipophilic or a hydrophilic amino acid or is deleted;

A³⁷ is a lipophilic amino acid or is deleted;

30 A³⁸ is a lipophilic or a hydrophilic amino acid or is deleted;

R¹ and R² are each independently selected from the group consisting of H, (C_{1-30}) alkyl, (C_{2-30}) alkenyl, phenyl- (C_{1-30}) alkyl, naphthyl (C_{1-30}) alkyl, hydroxy (C_{1-30}) alkyl, hydroxy (C_{2-30}) alkenyl, hydroxy-phenyl (C_{1-30}) alkyl or hydroxy-naphthyl (C_{1-30}) alkyl;

WO 99/57139 PCT/US99/09521

or one of R¹ or R² is COE¹ where E¹ is $(C_{1\cdot30})$ alkyl, $(C_{2\cdot3c})$ alkenyl, phenyl $(C_{1\cdot30})$ alkyl, naphthyl $(C_{1\cdot30})$ alkyl, hydroxy $(C_{1\cdot30})$ alkyl, hydroxy $(C_{2\cdot30})$ alkenyl, hydroxy-phenyl $(C_{1\cdot30})$ alkyl or hydroxy-naphthyl $(C_{1\cdot30})$ alkyl; and R³ is OH, NH₂, $(C_{1\cdot30})$ alkoxy or NH-Y-CH₂-Z, where Y is a $(C_{1\cdot30})$ hydrocarbon moiety and Z is CO₂H or CONH₂;

provided that the compound is not PTH(1-34)R 3 , PTH(1-35)R 3 , PTH(1-36)R 3 , PTH(1-37)R 3 , or PTH(1-38)R 3 .

- 8. A method of selectively binding a PTH2 receptor which comprises administering to a patient in need thereof an effective amount of an analogue according to claim 7 or a pharmaceutically-acceptable salt thereof.
- 9. An analogue according to claim 1 of formula (II), $(R^1R^2) A^1 A^2 A^3 A^4 A^5 A^6 A^7 A^8 A^{10} A^{11} A^{12} A^{13} A^{14} A^{15} A^{16} A^{17} A^{18} A^{19} A^{20} A^{21} A^{22} A^{23} A^{24} A^{25} A^{26} A^{27} A^{28} A^{30} A^{31} A^{32} A^{33} A^{34} A^{35} A^{36} A^{37} A^{38} R^3,$

(II)

or a pharmaceutically-acceptable salt thereof wherein

A1 is Ser. Ala, Dap, Thr, Aib or is deleted;

A² is Val, Leu, Ile, Phe, NIe, β-Nal, Aib, p-X-Phe, Acc, Cha, Met or is deleted;

A³ is Ser, Thr, Aib or is deleted;

A⁴ is Glu, Asp or is deleted;

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20 A⁵ is Leu, Val, Nle, Ile, Cha, β-Nal, Trp, Pal, Acc, Phe, p-X-Phe or is deleted;

A⁶ is Gln, a hydrophilic amino acid or is deleted;

 A^7 is Leu, Val, NIe, Ile, Cha, β -Nal, Trp, Pal, Acc, Phe, p-X-Phe, a lipophilic amino acid, or is deleted;

A⁸ is Met, Nva, Leu, Val, Ile, Cha, Acc, Nle, p-X-Phe, Phe, β -Nal, Bpa, a lipophilic amino acid or is deleted;

A9 is His, a hydrophilic amino acid or is deleted;

A¹⁰ is Asn, a hydrophilic amino acid or is deleted;

A¹¹ is Leu, Val, NIe, IIe, Cha, β-Nal, Trp, Pal, Acc, Phe, p-X-Phe, a hydrophilic amino acid or is deleted;

30 A¹² is Gly, Acc, Aib, or is deleted;

 A^{13} is Lys, Arg or HN-CH((CH₂)₀NH-R⁴)-C(O);

A14 is His or is deleted;

A¹⁵ is Leu, Val, Nle, Ile, Cha, β-Nal, Trp, Pal, Acc, Phe, p-X-Phe or is deleted;

A¹⁶ is Ser. Asn. Ala. Aib or is deleted;

35 A¹⁷ is Ser, Thr, Aib or is deleted;

A¹⁸ is Met, Nva, Leu, Val, Ile, Nle, p-X-Phe, Phe, β -Nal, Acc, Cha, Aib or is deleted; A¹⁹ is Glu, Aib or is deleted;

A²⁰ is Arg. Lys, HN-CH((CH₂),NH-R⁴)-C(O) or is deleted;

A²¹ is Val. Leu, Ile, Phe, Nle, β-Nal, Aib, p-X-Phe, Acc, Cha, Met or is deleted;

5 A²² is Acc, Aib, Glu or is deleted;

A²³ is Trp, Acc, Phe, p-X-Phe, Aib, β-Nal or Cha;

A²⁴ is Leu, Acc, Ile, Val, Phe, β-Nal, Nle, Aib, p-X-Phe or Cha;

 A^{25} is Arg. Lys or HN-CH((CH₂)₂NH-R⁴)-C(O);

 A^{26} is Arg, Lys or HN-CH((CH₂)₀NH-R⁴)-C(O);

10 A^{27} is Lys, Aib, Leu, hArg, Gln, Acc, Arg, Cha, Nle, Ile, Val, Phe, β -Nal, or p-X-Phe, where the Lys is optionally substituted on the ϵ -amino group by an acyl group;

A²⁸ is Leu, Acc, Cha, Ile, Val, Phe, Nle, β-Nal, Aib or p-X-Phe;

A²⁹ is Gln, Acc or Aib;

A³⁰ is Asp, Lys, Arg or is deleted;

15 A^{31} is Val, Leu, Nle, Acc, Cha, Phe, Ile, β -Nal Aib, p-X-Phe or is deleted;

A³² is His or is deleted;

A³³ is Asn or is deleted;

25

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A³⁴ is Phe. Tyr. Amp, Aib, β-Nal, Cha, Nle, Leu, Ile, Acc, p-X-Phe or is deleted;

A³⁵ is Val, Leu, Nle, Acc, Cha, Phe, Ile, β-Nal Aib, p-X-Phe or is deleted;

20 A³⁶ is Ala. Val. Aib. Acc. Nva. Abu or is deleted;

A³⁷ is Leu, Val, Nle, Ile, Cha, β-Nal, Trp, Pal, Acc, Phe, p-X-Phe, a lipophilic amino acid, or is deleted;

A³⁸ is Gly, Acc, Aib, or is deleted;

where X for each occurrence is independently selected from the group consisting of OH, a halo and CH₃;

R¹ and R² are each independently selected from the group consisting of H, (C_{1-30}) alkyl, (C_{2-30}) alkenyl, phenyl- (C_{1-30}) alkyl, naphthyl (C_{1-30}) alkyl, hydroxy (C_{1-30}) alkyl, hydroxy (C_{1-30}) alkyl, hydroxy-phenyl (C_{1-30}) alkyl; or hydroxy-naphthyl (C_{1-30}) alkyl;

or one of R¹ or R² is COE¹ where E¹ is (C_{1-30}) alkyl, (C_{2-30}) alkenyl, phenyl (C_{1-30}) alkyl, naphthyl (C_{1-30}) alkyl, hydroxy (C_{1-30}) alkyl, hydroxy (C_{2-30}) alkenyl, hydroxy-phenyl (C_{1-30}) alkyl or hydroxy-naphthyl (C_{1-30}) alkyl;

 R^3 is OH, NH_2 , $(C_{1.30})$ alkoxy or $NH-Y-CH_2-Z$, where Y is a $(C_{1.30})$ hydrocarbon moiety and Z is CO_2H or $CONH_2$;

n for each occurrence is independently an integer from 1 to 5; and

5

WO 99/57139 PCT/US99/09521 36

> R4 for each occurrence is independently (C1-C30)alkyl, (C1-C30)acyl or - $C((NH)(NH_2));$

provided that the compound is not PTH(1-34)R3, PTH(1-35)R3, PTH(1-36)R3, PTH(1-36)R 37)R³, or PTH(1-38)R³.

> A compound of the formula (III), 10.

 $(R^{1}R^{2})-A^{1}-A^{2}-A^{3}-A^{4}-A^{5}-A^{6}-A^{7}-A^{8}-A^{9}-A^{10}-A^{11}-A^{12}-A^{13}-A^{14}-A^{15}-A^{16}-A^{17}-A^{18}-A^{19}-A^{20}-A^{21}-A^{22}-A^{23}-A^{10}-A^{11}-A^{12}-A^{12}-A^{13}-A^{14}-A^{15}-A^{16}-A^{17}-A^{18}-A^{19}-A^{20}-A^{21}-A^{22}-A^{23}-A^{21}-A^{22}-A^{23}-A^{21}-A^{22}-A^{23}-A^{$ $A^{24} - A^{25} - A^{26} - A^{27} - A^{28} - A^{29} - A^{30} - A^{31} - A^{32} - A^{33} - A^{34} - A^{35} - A^{36} - A^{37} - A^{38} - R^3$.

(III)

or a pharmaceutically-acceptable salt thereof wherein

10 A¹ is Ser, Ala, Dap, Thr, Aib or is deleted;

A² is Val, Leu, Ile, Phe, Nle, β-Nal, Aib, p-X-Phe, Acc, Cha, Met or is deleted;

A³ is Ser, Thr, Aib or is deleted;

A4 is Glu, Asp or is deleted;

A⁵ is Leu, Val, Nie, Ile, Cha, β-Nal, Trp, Pal, Acc, Phe, p-X-Phe or is deleted;

15 A⁶ is Gln, a hydrophilic amino acid or is deleted;

 A^7 is Leu, Val, NIe, IIe, Cha, β -Nal, Trp, Pal, Acc, Phe, p-X-Phe, a lipophilic amino acid, or is deleted;

 A^8 is Met, Nva, Leu, Val, Ile, Cha, Acc, Nle, p-X-Phe, Phe, β -Nal, Bpa, a lipophilic amino acid or is deleted;

20 A⁹ is His, a hydrophilic amino acid or is deleted;

A¹⁰ is Asn, a hydrophilic amino acid or is deleted;

A¹¹ is Leu, Val, NIe, IIe, Cha, β-Nal, Trp, Pal, Acc, Phe, p-X-Phe, a hydrophilic amino acid or is deleted;

A¹² is Gly, Acc, Aib, or is deleted;

A¹³ is Lys. Arg or HN-CH((CH₂)₀NH-R⁴)-C(O); 25

A¹⁴ is His or is deleted;

A¹⁵ is Leu, Val, Nle, Ile, Cha, β-Nal, Trp, Pal, Acc, Phe, p-X-Phe or is deleted;

A¹⁶ is Ser, Asn, Ala, Aib or is deleted;

A¹⁷ is Ser, Thr, Aib or is deleted;

A¹⁸ is Met, Nva, Leu, Val; Ile, Nle, p-X-Phe, Phe, β-Nal, Acc, Cha, Aib or is deleted; 30

A¹⁹ is Glu, Aib or is deleted;

A²⁰ is Arg, Lys, HN-CH((CH₂)₀NH-R⁴)-C(O) or is deleted;

A²¹ is Val. Leu. Ile. Phe. Nie. β-Nai, Aib, p-X-Phe, Acc, Cha, Met or is deleted;

A²² is Acc. Aib. Glu or is deleted;

35 A^{23} is Trp. Acc, Phe, p-X-Phe, Aib, β -Nal or Cha;

A24 is Leu, Acc, Ile, Val, Phe, B-Nal, Nle, Aib, p-X-Phe or Cha;

 A^{25} is Arg, Lys or HN-CH((CH₂)₀NH-R⁴)-C(O);

 A^{26} is Arg, Lys or HN-CH((CH₂)_nNH-R⁴)-C(O);

A²⁷ is Lys, Aib, Leu, hArg, Gln, Acc, Arg, Cha, Nle, Ile, Val, Phe, β-Nal, or p-X-Phe,

5 where the Lys is optionally substituted on the ϵ -amino group by an acyl group;

A²⁸ is Leu, Acc, Cha, Ile, Val, Phe, Nle, β-Nal, Aib or p-X-Phe;

A²⁹ is Gln. Acc or Aib;

A³⁰ is Asp. Lys. Arg or is deleted;

 A^{31} is Val. Leu, NIe, Acc, Cha, Phe, IIe, β -Nal Aib, p-X-Phe or is deleted;

10 A³² is His or is deleted;

25

30

A³³ is Asn or is deleted;

A³⁴ is Phe, Tyr, Amp, Aib, β-Nal, Cha, Nle, Leu, Ile, Acc, p-X-Phe or is deleted;

A³⁵ is Val, Leu, Nle, Acc, Cha, Phe, Ile, β-Nal Aib, p-X-Phe or is deleted;

A³⁶ is Ala, Val, Aib, Acc, Nva, Abu or is deleted;

15 A³⁷ is Leu, Val, Nle, Ile, Cha, β-Nal, Trp, Pal, Acc, Phe, p-X-Phe, a lipophilic amino acid, or is deleted;

A³⁸ is Gly, Acc, Aib, or is deleted;

where X for each occurrence is independently selected from the group consisting of OH, a halo and CH₃;

20 R¹ and R² are each independently selected from the group consisting of H, $(C_{1-30}) \text{alkyl}, \quad (C_{2-30}) \text{alkenyl}, \quad \text{phenyl-}(C_{1-30}) \text{alkyl}, \quad \text{naphthyl}(C_{1-30}) \text{alkyl}, \\ \text{hydroxy}(C_{1-30}) \text{alkyl}, \quad \text{hydroxy}(C_{2-30}) \text{alkenyl}, \quad \text{hydroxy-phenyl}(C_{1-30}) \text{alkyl} \quad \text{or} \\ \text{hydroxy-naphthyl}(C_{1-30}) \text{alkyl};$

or one of R¹ or R² is COE¹ where E¹ is (C_{1-30}) alkyl, (C_{2-30}) alkenyl, phenyl (C_{1-30}) alkyl, naphthyl (C_{1-30}) alkyl, hydroxy (C_{1-30}) alkyl, hydroxy (C_{2-30}) alkenyl, hydroxy-phenyl (C_{1-30}) alkyl or hydroxy-naphthyl (C_{1-30}) alkyl;

 R^3 is OH, NH₂, (C₁₋₃₀)alkoxy or NH-Y-CH₂-Z, where Y is a (C₁₋₃₀) hydrocarbon moiety and Z is CO₂H or CONH₂;

n for each occurrence is independently an integer from 1 to 5; and

R⁴ for each occurrence is independently (C_1-C_{30}) alkyl, (C_1-C_{30}) acyl or - $C((NH)(NH_2))$;

provided that when A⁸ is not a lipophilic D-amino acid or is not deleted then at least one of A⁶, A⁷, A⁹, A¹⁰, A¹¹ and A¹² is a D-amino acid or at least one of A⁶, A⁷, A⁹, A¹⁰, A¹¹, A¹², A¹³, A¹⁴, A¹⁵, A¹⁶, A¹⁷, A¹⁸, A¹⁹, A²⁰, A²¹ and A²² is deleted;

and further provided that when the compound contains a D-amino acid then A³⁶ is deleted.

- 11. A compound according to claim 10 wherein said compound is [D-Nie⁸, Nie¹⁸, Tyr³⁴]hPTH(1-34)NH₂,
- 5 [D-Nie⁸]hPTH(1-34)NH₂,

[D-Leu⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂,

[D-Cha⁶, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂,

[D-Phe⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂,

[D-Nal8, Nie18, Tyr34]hPTH(1-34)NH₂,

10 [D-Abu⁸, Nie¹⁸, Tyr³⁴]hPTH(1-34)NH₂,

[D-Met⁸]hPTH(1-34)NH₂,

[Cha^{7, 11}, D-Met⁸]hPTH(1-34)NH₂,

[D-lie⁸]hPTH(1-34)NH₂,

[Cha^{7, 11}, D-Ile⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂,

- 15 [D-lle⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂,
 - [D-Leu⁸]hPTH(1-34)NH₂,

[Cha^{7,11}, D-Leu⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂,

[D-Val8]hPTH(1-34)NH₂,

[Cha^{7, 11}, D-Val⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂,

- 20 [D-Val⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂,
 - [D-Cha8]hPTH(1-34)NH₂,

[Cha^{7,11}, D-Cha⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂,

 $[D-Ala^8]hPTH(1-34)NH_2$

[Cha^{7, 11}, D-Ala⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂,

- 25 [D-Ala⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂,
 - [D-Phe⁸]hPTH(1-34)NH₂,

[Cha^{7,11}, D-Phe⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂,

[D-Nal⁸]hPTH(1-34)NH₂,

[D-Trp⁸]hPTH(1-34)NH₂,

- 30 [Cha^{7,11}, D-Trp⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂,
 - [D-Trp8, Nle18, Tyr34]hPTH(1-34)NH₂,

[D-Abu⁸]hPTH(1-34)NH₂,

[Cha^{7,11}, D-Abu⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂,

[D-Nie⁸, Nie¹⁸]hPTH(1-34)NH₂,

35 [des-Met⁸]hPTH(1-34)NH₂,

[Cha^{7.11}, des-Met⁸]hPTH(1-34)NH₂,
[Cha^{7.11}, des-Met⁸, des-Met¹⁸, Tyr³⁴]hPTH(1-34)NH₂,
[des-Met⁸, des-Met¹⁸]hPTH(1-34)NH₂,
[Cha^{7.11}, des-Met⁸, des-Met¹⁸]hPTH(1-34)NH₂,
[des-Met⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂,
[des-Met¹⁸]hPTH(1-34)NH₂,
[Cha^{7.11}, des-Met¹⁸]hPTH(1-34)NH₂,
[Cha^{7.11}, des-Met¹⁸, Tyr³⁴]hPTH(1-34)NH₂,
[D-Nle⁸, des-Met¹⁸, Tyr³⁴]hPTH(1-34)NH₂,
[des-Glu⁶, Nle^{8,18}, Tyr³⁴]hPTH(1-34)NH₂,
[des-Leu⁷, Nle^{8,18}, Tyr³⁴]hPTH(1-34)NH₂,
[des-His⁹, Nle^{8,18}, Tyr³⁴]hPTH(1-34)NH₂,

[des-Gly¹², Nle^{8,18}, Tyr³⁴]hPTH(1-34)NH₂, [des-Lys¹³, Nle^{8,18}, Tyr³⁴]hPTH(1-34)NH₂, [des-His¹⁴, Nle^{8,18}, Tyr³⁴]hPTH(1-34)NH₂, [des-Leu¹⁵, Nle^{8,18}, Tyr³⁴]hPTH(1-34)NH₂, [des-Asn¹⁶, Nle^{8,18}, Tyr³⁴]hPTH(1-34)NH₂,

[des-Leu¹¹, Nle^{8,18}, Tyr³⁴]hPTH(1-34)NH₂,

- 20 [des-Ser¹⁷, Nle^{8,18}, Tyr³⁴]hPTH(1-34)NH₂, [des-Glu¹⁹, Nle^{8,18}, Tyr³⁴]hPTH(1-34)NH₂, [des-Arg²⁰, Nle^{8,18}, Tyr³⁴]hPTH(1-34)NH₂, [des-Val²¹, Nle^{8,18}, Tyr³⁴]hPTH(1-34)NH₂, [des-Glu²², Nle ^{8,18}, Tyr³⁴]hPTH(1-34)NH₂,
- 25 [des-Glu⁶, Cha^{7,11}, Nle^{8,18}, Tyr³⁴]hPTH(1-34)NH₂,
 [des-Leu⁷, Nle^{8,18}, Cha¹¹, Tyr³⁴]hPTH(1-34)NH₂,
 [Cha^{7,11}, des-His⁹, Nle^{8,18}, Tyr³⁴]hPTH(1-34)NH₂,
 [des-Glu⁶, Cha^{7,11}, D-Nle⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂,
 [des-Leu⁷, D-Nle⁸, Cha¹¹, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂,
- [Cha^{7,11}, D-Nle⁸, des-His⁹, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂,
 [Cha^{7,11}, D-Nle⁸, Nle¹⁸, Tyr³⁴]hPTH(1-31)NH₂,
 [Cha^{7,11}, des-Met⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂,
 [Cha^{7,11}, D-Nle⁸, des-Met¹⁸, Tyr³⁴]hPTH(1-34)NH₂,
 [Cha^{7,11}, des-Met⁸, des-His⁹, des-Asn¹⁰]hPTH(1-34)NH₂,
- 35 [Cha^{7,11}, des-Ser¹⁷, des-Met¹⁸, des-Glu¹⁹]hPTH(1-34)NH₂,

[D-Met⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂,

[D-Met⁸, Tyr³⁴]hPTH(1-34)NH₂,

[D-Bpa8, Tyr34]hPTH(1-34)NH₂,

[D-Nie⁸, Nie¹⁸, Tyr³⁴]hPTH(7-34)NH₂,

5 [D-Nle⁸, Nle¹⁸]hPTH(7-34)NH₂ or

[D-Met⁸]hPTH(7-34)NH₂.

12. A compound according to claim 11 wherein said compound is

[Cha^{7,11}, des-Met⁸, Nle¹⁸, Tyr³⁴]hPTH-(1-34)NH₂,

[Cha^{7,11}, D-Nle⁸, des-Met¹⁸, Tyr³⁴]hPTH-(1-34)NH₂,

10 [Cha^{7,11}, D-Nle⁸, Nle¹⁸, Tyr³⁴]hPTH-(1-34)NH₂,

[D-Nle⁸, Nle¹⁸, Tyr³⁴]hPTH(1-34)NH₂ or [D-Bpa⁸, Tyr³⁴]hPTH(1-34)NH₂.

13. A PTHrP analogue of formula (IV) that selectively binds to the PTH2 receptor,

 $(R^1R^2) - A^1 - A^2 - A^3 - A^4 - A^5 - A^6 - A^7 - A^8 - A^9 - A^{10} - A^{11} - A^{12} - A^{13} - A^{14} - A^{15} - A^{16} - A^{17} - A^{18} - A^{19} - A^{20} - A^{21} - A^{22} - A^{23} - A^{20} -$

15 A^{24} - A^{25} - A^{26} - A^{27} - A^{28} - A^{29} - A^{30} - A^{31} - A^{32} - A^{33} - A^{34} - A^{35} - A^{36} - A^{37} - A^{38} - R^{3} ,

(IV)

or a pharmaceutically acceptable salt thereof, wherein

A1 is Ala, Ser, Dap, Thr, Aib or is deleted;

A² is Val or is deleted;

20 A³ is Ser, Aib, Thr or is deleted;

A4 is Glu, Asp or is deleted;

A⁵ is His, Ile, Acc, Val, Nle, Phe, Leu, p-X-Phe, β-Nal, Aib, Cha or is deleted;

A⁶ is Gln, a hydrophilic amino acid or is deleted;

A⁷ is Leu, Val, Cha, Nle, β-Nal, Trp, Pal, Acc, Phe, p-X-Phe, Aib, a lipophilic amino

25 acid or is deleted;

A⁸ is Leu, Met, Acc, Cha, Aib, Nle, Phe, Ile, Val, β -Nal, p-X-Phe, a lipophilic amino acid or is deleted;

A9 is His, a hydrophilic amino acid or is deleted;

A¹⁰ is Asp. Asp. a hydrophilic amino acid or is deleted;

30 A¹¹ is Lys, Arg, Leu, Cha, Aib, p-X-Phe, Ile, Val, Nle, Acc, Phe, β-Nal, HN-CH((CH₂)_nNH-R⁴)-C(O), a lipophilic D-amino acid, a hydrophilic amino acid or is deleted;

A12 is Gly, Acc, Aib or is deleted;

A¹³ is Lys. Arg. HN-CH((CH₂)₂NH-R⁴)-C(O) or is deleted;

35 A¹⁴ is Ser, His or is deleted;

A¹⁵ is Ile, Acc, Cha, Leu, Phe, Nle, *B*-Nal, Trp, p-X-Phe, Val, Aib or is deleted;

A¹⁶ is Gln. Aib or is deleted;

A¹⁷ is Asp, Aib or is deleted;

A¹⁸ is Leu, Aib, Acc, Cha, Phe, Ile, Nle, β-Nal, Val, p-X-Phe or is deleted;

5 A¹⁹ is Arg, Lys, Aib, HN-CH((CH₂), NH-R⁴)-C(O) or is deleted;

A²⁰ is Arg, Lys, HN-CH((CH₂)_nNH-R⁴)-C(O) or is deleted;

A²¹ is Arg, Lys, HN-CH((CH₂)_nNH-R⁴)-C(O) or is deleted;

A²² is Phe, Glu, Aib, Acc, p-X-Phe, β-Nal, Val, Leu, Ile, Nle or Cha;

A²³ is Phe, Leu, Lys, Acc, Cha, β-Nal, Aib, Nle, Ile, p-X-Phe, Val or Trp;

10 A^{24} is Leu, Lys, Acc, Nle, Ile, Val, Phe, β -Nal, Aib, p-X-Phe, Arg or Cha;

A²⁵ is His, Lys, Aib, Acc, Arg or Glu;

A²⁶ is His, Aib, Acc, Arg or Lys;

A²⁷ is Leu, Lys, Acc, Arg, Ile, Val, Phe, Aib, Nle, β-Nal, p-X-Phe or Cha;

A²⁶ is IIe, Leu, Lys, Acc, Cha, Val, Phe, p-X-Phe, NIe, β-Nal, Aib or is deleted;

15 A²⁹ is Ala, Glu, Acc, Aib or is deleted;

A³⁰ is Glu, Leu, Nle, Cha, Aib, Acc, Lys, Arg or is deleted;

A³¹ is Ile, Leu, Cha, Lys, Acc, Phe, Val, Nle, β-Nal, Arg or is deleted;

A³² is His or is deleted;

25

30

A³³ is Thr. Ser or is deleted;

20 A^{34} is Ala, Phe, Tyr, Cha, Val, IIe, Leu, NIe, β -Nal, Aib, Acc or is deleted;

A³⁵ is Glu, Asp or is deleted;

A³⁶ is Ile, Acc, Cha, Leu, Phe, Nle, β-Nal, Trp, p-X-Phe, Val, Aib or is deleted;

A³⁷ is Arg, Lys, HN-CH((CH₂)_nNH-R⁴)-C(O) or is deleted;

A³⁸ is Ala, Phe, Tyr, Cha, Val, Ile, Leu, Nle, *β*-Nal, Aib, Acc or is deleted;

R¹ and R² are each independently selected from the group consisting of H, (C_{1-30}) alkyl, (C_{2-30}) alkenyl, phenyl- (C_{1-30}) alkyl, naphthyl (C_{1-30}) alkyl, hydroxy (C_{1-30}) alkyl, hydroxy (C_{1-30}) alkyl, hydroxy-phenyl (C_{1-30}) alkyl or hydroxy-naphthyl (C_{1-30}) alkyl;

or one of R¹ or R² is COE¹ where E¹ is $(C_{1.30})$ alkyl, $(C_{2.30})$ alkenyl, phenyl $(C_{1.30})$ alkyl, naphthyl $(C_{1.30})$ alkyl, hydroxy $(C_{1.30})$ alkyl, hydroxy $(C_{1.30})$ alkyl, hydroxy-phenyl $(C_{1.30})$ alkyl or hydroxy-naphthyl $(C_{1.30})$ alkyl;

 R^3 is OH, NH_2 , (C_{1-30}) alkoxy or $NH-Y-CH_2-Z$, where Y is a (C_{1-30}) hydrocarbon moiety and Z is CO_2H or $CONH_2$;

n for each occurrence is independently an integer from 1 to 5; and

 R^4 for each occurrence is independently (C_1-C_{30}) alkyl, (C_1-C_{30}) acyl or - $C((NH)(NH_2))$;

provided that the compound is not PTHrP(1-34)R³, PTHrP(1-35)R³, PTHrP(1-36)R³, PTHrP(1-37)R³ or PTHrP(1-38)R³,

5 and further provided that the compound is not [lle⁵, Trp²³]PTHrP(1-36) or [Trp²³]PTHrP(1-36).

14. A compound of formula (V),

 $(\mathsf{R}^1\mathsf{R}^2) - \mathsf{A}^1 - \mathsf{A}^2 - \mathsf{A}^3 - \mathsf{A}^4 - \mathsf{A}^5 - \mathsf{A}^6 - \mathsf{A}^7 - \mathsf{A}^8 - \mathsf{A}^9 - \mathsf{A}^{10} - \mathsf{A}^{11} - \mathsf{A}^{12} - \mathsf{A}^{13} - \mathsf{A}^{14} - \mathsf{A}^{15} - \mathsf{A}^{16} - \mathsf{A}^{17} - \mathsf{A}^{18} - \mathsf{A}^{19} - \mathsf{A}^{20} - \mathsf{A}^{21} - \mathsf{A}^{22} - \mathsf{A}^{23} - \mathsf{A}^{24} - \mathsf{A}^{25} - \mathsf{A}^{26} - \mathsf{A}^{27} - \mathsf{A}^{28} - \mathsf{A}^{29} - \mathsf{A}^{30} - \mathsf{A}^{31} - \mathsf{A}^{32} - \mathsf{A}^{33} - \mathsf{A}^{34} - \mathsf{A}^{35} - \mathsf{A}^{36} - \mathsf{A}^{37} - \mathsf{A}^{38} - \mathsf{R}^3 \ ,$

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or a pharmaceutically acceptable salt thereof, wherein

A1 is Ala, Ser, Dap, Thr, Aib or is deleted;

A² is Val or is deleted;

A³ is Ser, Aib, Thr or is deleted;

15 A4 is Glu, Asp or is deleted;

A⁵ is His, Ile, Acc, Val, Nle, Phe, Leu, p-X-Phe, β-Nal, Aib, Cha or is deleted;

A⁶ is Gln, a hydrophilic amino acid or is deleted;

 A^7 is Leu, Val, Cha, Nle, β -Nal, Trp, Pal, Acc, Phe, p-X-Phe, Aib, a lipophilic amino acid or is deleted;

A⁸ is Leu, Met, Acc, Cha, Aib, Nle, Phe, Ile, Val, β-Nal, p-X-Phe, a lipophilic amino acid or is deleted;

A⁹ is His, a hydrophilic amino acid or is deleted:

A¹⁰ is Asp. Asn. a hydrophilic amino acid or is deleted;

A¹¹ is Lys, Arg, Leu, Cha, Aib, p-X-Phe, Ile, Val, Nle, Acc, Phe, β-Nal, HN-

25 CH((CH₂)_nNH-R⁴)-C(O), a lipophilic D-amino acid, a hydrophilic amino acid or is deleted:

A¹² is Gly, Acc, Aib or is deleted;

A¹³ is Lys, Arg, HN-CH((CH₂)_cNH-R⁴)-C(O) or is deleted;

A¹⁴ is Ser, His or is deleted;

30 A^{15} is IIe, Acc, Cha, Leu, Phe, NIe, β -Nal, Trp, p-X-Phe, Val, Aib or is deleted;

A¹⁶ is Gln. Aib or is deleted;

A¹⁷ is Asp, Aib or is deleted;

A¹⁸ is Leu, Aib, Acc, Cha, Phe, Ile, Nle, β-Nal, Val, p-X-Phe or is deleted;

A¹⁹ is Arg, Lys, Aib, HN-CH((CH₂), NH-R⁴)-C(O) or is deleted;

35 A^{20} is Arg, Lys, HN-CH((CH₂)₀NH-R⁴)-C(O) or is deleted;

WO 99/57139 43 PCT/US99/09521

A²¹ is Arg. Lys, HN-CH((CH₂)_nNH-R⁴)-C(O) or is deleted;

A²² is Phe, Glu, Aib, Acc, p-X-Phe, β-Nal, Val, Leu, Ile, Nle or Cha;

 A^{23} is Phe, Leu, Lys, Acc, Cha, β -Nal, Aib, Nie, Ile, p-X-Phe, Val or Trp;

A²⁴ is Leu, Lys, Acc, Nle, Ile, Val, Phe, β-Nal, Aib, p-X-Phe, Arg or Cha;

5 A²⁵ is His, Lys, Aib, Acc, Arg or Glu;

A²⁶ is His, Aib, Acc, Arg or Lys;

A²⁷ is Leu, Lys, Acc, Arg, Ile, Val, Phe, Aib, Nle, β-Nal, p-X-Phe or Cha;

A²⁸ is IIe. Leu. Lvs. Acc. Cha, Val. Phe, p-X-Phe, NIe, β-Nal, Aib or is deleted;

A²⁹ is Ala, Glu, Acc, Aib or is deleted;

10 A³⁰ is Glu, Leu, Nle, Cha, Aib, Acc, Lys, Arg or is deleted;

 A^{31} is IIe, Leu, Cha, Lys, Acc, Phe, Val, NIe, β -Nal, Arg or is deleted;

A³² is His or is deleted;

A³³ is Thr, Ser or is deleted;

A³⁴ is Ala, Phe, Tyr, Cha, Val, Ile, Leu, Nle, β-Nal, Aib, Acc or is deleted;

15 A³⁵ is Glu, Asp or is deleted;

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A³⁶ is IIe, Acc, Cha, Leu, Phe, NIe, β-Nal, Trp, p-X-Phe, Val, Aib or is deleted;

A³⁷ is Arg, Lys, HN-CH((CH₂),NH-R⁴)-C(O) or is deleted;

A³⁸ is Ala, Phe, Tyr, Cha, Val, Ile, Leu, Nle, β-Nal, Aib, Acc or is deleted;

 R^1 and R^2 are each independently selected from the group consisting of H, (C_{1-30}) alkyl, (C_{2-30}) alkenyl, phenyl- (C_{1-30}) alkyl, naphthyl (C_{1-30}) alkyl, hydroxy (C_{1-30}) alkyl, hydroxy-phenyl (C_{1-30}) alkyl, or hydroxy-naphthyl (C_{1-30}) alkyl;

or one of R^1 or R^2 is COE^1 where E^1 is (C_{1-30}) alkyl, (C_{2-30}) alkenyl, phenyl(C_{1-30})alkyl, naphthyl(C_{1-30})alkyl, hydroxy(C_{1-30})alkyl, hydroxy-phenyl(C_{1-30})alkyl or hydroxy-naphthyl(C_{1-30})alkyl;

 R^3 is OH, NH_2 , (C_{1-30}) alkoxy or $NH-Y-CH_2-Z$, where Y is a (C_{1-30}) hydrocarbon moiety and Z is CO_2H or $CONH_2$;

n for each occurrence is independently an integer from 1 to 5; and R^4 for each occurrence is independently (C_1-C_{30}) alkyl, (C_1-C_{30}) acyl or $-C((NH)(NH_2))$;

provided that when A⁸ is not a lipophilic D-amino acid or is not deleted then at least one of A⁶, A⁷, A⁹, A¹⁰, A¹¹ and A¹² is a D-amino acid or at least one of A⁶, A⁷, A⁹, A¹⁰, A¹¹, A¹², A¹³, A¹⁴, A¹⁵, A¹⁶, A¹⁷, A¹⁸, A¹⁹, A²⁰, A²¹ and A²² is deleted.

15. A compound according to claim 14 wherein said compound is [Ile⁵, D-Leu⁸]hPTHrP(1-34)NH₂,

[Ile⁵, D-Leu⁸, Trp²³]hPTHrP(1-34)NH₂,
[Ile⁵, des-Leu⁸]hPTHrP(1-34)NH₂,
[Ile⁵, des-Leu⁸]hPTHrP(1-34)NH₂,
[des-Leu⁸, Trp²³]hPTHrP(1-34)NH₂,
[Ile⁵, des-Leu¹⁸]hPTHrP(1-34)NH₂,
[Ile⁵, des-Leu¹⁸, Trp²³]hPTHrP(1-34)NH₂,
[des-Leu¹⁸, Trp²³]hPTHrP(1-34)NH₂,
[Ile⁵, D-Leu⁸, Glu^{22,25}, Leu^{23,28,31}, Lys^{26,30}, Aib²⁹]hPTHrP(1-34)NH₂,
[Ile⁵, D-Leu⁸, Glu^{22,25}, Trp²³, Lys^{26,30}, Leu^{28,31}, Aib²⁹]hPTHrP(1-34)NH₂,
[Ile⁵, D-Leu⁸, Glu^{22,25,29}, Leu^{23,28,31}, Lys^{26,30}]hPTHrP(1-34)NH₂,
[Ile⁵, D-Leu⁸, Glu^{22,25,29}, Trp²³, Lys^{26,30}, Leu^{28,31}]hPTHrP(1-34)NH₂ or
[D-Leu⁸, Trp²³]hPTHrP(7-34)NH₂.

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- 16. A method of selectively binding the PTH2 receptor which comprises administering to a patient in need thereof an analogue according to claim 9 or a pharmaceutically acceptable salt thereof.
- 17. A method of selectively binding the PTH2 receptor which comprises administering to a patient in need thereof a compound according to claim 10 or a pharmaceutically acceptable salt thereof.
- 18. A method of selectively binding the PTH2 receptor which comprises 20 administering to a patient in need thereof a compound according to claim 11 or a pharmaceutically acceptable salt thereof.
 - 19. A method of selectively binding a PTH2 receptor which comprises administering to a patient in need thereof a compound according to claim 12 or a pharmaceutically acceptable salt thereof.
 - 20. A method of selectively binding a PTH2 receptor which comprises administering to a patient in need thereof an analogue according to claim 13 or a pharmaceutically acceptable salt thereof.
 - 21. A method of selectively binding a PTH2 receptor which comprises administering to a patient in need thereof a compound according to claim 14 or a pharmaceutically acceptable salt thereof.
 - 22. A method of selectively binding a PTH2 receptor which comprises administering to a patient in need thereof a compound according to claim 15 or a pharmaceutically acceptable salt thereof.

- 23. A pharmaceutical composition comprising an analogue according to claim 9 or a pharmaceutically acceptable salt thereof and a pharmaceutically acceptable carrier.
- 24. A pharmaceutical composition comprising a compound according
 to claim 10 or a pharmaceutically acceptable salt thereof and a pharmaceutically acceptable carrier.
 - 25. A pharmaceutical composition comprising a compound according to claim 11 or a pharmaceutically acceptable salt thereof and a pharmaceutically acceptable carrier.
- 26. A pharmaceutical composition comprising a compound according to claim 12 or a pharmaceutically acceptable salt thereof and a pharmaceutically acceptable carrier.
 - 27. A pharmaceutical composition comprising an analogue according to claim 13 or a pharmaceutically acceptable salt thereof and a pharmaceutically acceptable carrier.

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- 28. A pharmaceutical composition comprising a compound according to claim 14 or a pharmaceutically acceptable salt thereof and a pharmaceutically acceptable carrier.
- 29. A pharmaceutical composition comprising a compound according to claim 15 or a pharmaceutically acceptable salt thereof and a pharmaceutically acceptable carrier.
 - 30. A method of treating a medical disorder that results from altered or excessive action of the PTH2 receptor, which comprises administering to a patient in need thereof an effective amount of an analogue according to claim 7, sufficient to inhibit the activation of the PTH2 receptor of said patient.
 - 31. A method of treating a medical disorder that results from altered or excessive action of the PTH2 receptor, which comprises administering to a patient in need thereof an effective amount of an analogue according to claim 9, sufficient to inhibit the activation of the PTH2 receptor of said patient.
 - 32. A method of treating a medical disorder that results from altered or excessive action of the PTH2 receptor, which comprises administering to a patient in need thereof an effective amount of a compound according to claim 10, sufficient to inhibit the activation of the PTH2 receptor of said patient.
- 33. A method of treating a medical disorder that results from altered or
 excessive action of the PTH2 receptor, which comprises administering to a patient in

need thereof an effective amount of a compound according to claim 11, sufficient to inhibit the activation of the PTH2 receptor of said patient.

- 34. A method of treating a medical disorder that results from altered or excessive action of the PTH2 receptor, which comprises administering to a patient in need thereof an effective amount of a compound according to claim 12, sufficient to inhibit the activation of the PTH2 receptor of said patient.
- 35. A method of treating a medical disorder that results from altered or excessive action of the PTH2 receptor, which comprises administering to a patient in need thereof an effective amount of an analogue according to claim 13, sufficient to inhibit the activation of the PTH2 receptor of said patient.

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- 36. A method of treating a medical disorder that results from altered or excessive action of the PTH2 receptor, which comprises administering to a patient in need thereof an effective amount of a compound according to claim 14, sufficient to inhibit the activation of the PTH2 receptor of said patient.
- 37. A method of treating a medical disorder that results from altered or excessive action of the PTH2 receptor, which comprises administering to a patient in need thereof an effective amount of a compound according to claim 15, sufficient to inhibit the activation of the PTH2 receptor of said patient.
- 38. A method according to claim 30 wherein said medical disorder is abnormal CNS functions, abnormal pancreatic functions, divergence from normal mineral metabolism and homeostasis, male infertility, abnormal blood pressure or a hypothalmic disease.
- 39. A method according to claim 31 wherein said medical disorder is abnormal CNS functions, abnormal pancreatic functions, divergence from normal mineral metabolism and homeostasis, male infertility, abnormal blood pressure or a hypothalmic disease.
- 40. A method according to claim 32 wherein said medical disorder is abnormal CNS functions, abnormal pancreatic functions, divergence from normal mineral metabolism and homeostasis, male infertility, abnormal blood pressure or a hypothalmic disease.
- 41. A method according to claim 33 wherein said medical disorder is abnormal CNS functions, abnormal pancreatic functions, divergence from normal mineral metabolism and homeostasis, male infertility, abnormal blood pressure or a hypothalmic disease.

- 42. A method according to claim 34 wherein said medical disorder is abnormal CNS functions, abnormal pancreatic functions, divergence from normal mineral metabolism and homeostasis, male infertility, abnormal blood pressure or a hypothalmic disease.
- 43. A method according to claim 35 wherein said medical disorder is abnormal CNS functions, abnormal pancreatic functions, divergence from normal mineral metabolism and homeostasis, male infertility, abnormal blood pressure or a hypothalmic disease.

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- 44. A method according to claim 36 wherein said medical disorder is abnormal CNS functions, abnormal pancreatic functions, divergence from normal mineral metabolism and homeostasis, male infertility, abnormal blood pressure or a hypothalmic disease.
- 45. A method according to claim 37 wherein said medical disorder is abnormal CNS functions, abnormal pancreatic functions, divergence from normal mineral metabolism and homeostasis, male infertility, abnormal blood pressure or a hypothalmic disease.
- 46. A method of treating a medical disorder that results from altered or excessive action of the PTH2 receptor, which comprises administering to a patient in need thereof an effective amount of a PTH analogue or a truncated PTH analogue or a pharmaceutically acceptable salt thereof according to claim 1, sufficient to inhibit the activation of the PTH2 receptor of said patient.
- 47. A method according to claim 46 wherein said medical disorder is abnormal CNS functions, abnormal pancreatic functions, divergence from normal mineral metabolism and homeostasis, male infertility, abnormal blood pressure or a hypothalmic disease.